

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)

ADOOT THE FATILITY (pieuse ensure un inju	illiution below is	s com	unieu n	Title medici	ii recorus)			
3. Patient's initials	first last							
4. Sex (circle)	MALE		FEMALE					
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)	UPPER	L	OWER					
8. Haematemesis <u>or</u> coffee-ground vomitus (circle)	YES			NO	Also circle YES if presence of blood in nasoga aspirate			d in nasogastric
9. Melaena <u>or</u> fresh blood per rectum (circle)	YES		NO Also circle YES if occult or gr rectal examination		or gross b	gross blood present on		
10. Suspected variceal bleed? (circle)	YES			NO				
11. Systolic blood pressure	Most recent measurement prior to randomisation mmHg				on			
12. Heart rate	Most recent meas			cent measurer	rement prior to randomisation			
13. Signs of shock present? (circle)	YES		NO		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)	YES		NO		Clinical judgement after considering history, signs and symptoms			
15. Major co-morbidities? (circle all that apply)	CARDIOVASCULAR	RESP	IRATORY	Liver	RENAL	MALI	GNANCY	OTHER MAJOR CO-MORBIDITY
16. On anti-coagulant therapy? (circle)	YES			NO	UNKNOWN	J		
17. Emergency admission? (circle)	YES	YES NO If patient already hospitalised, cit			alised, cir	cle '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)	YES				do not r	NO do not randomise, record on screening log			
19. Consent for entry obtained from (circle)	WAIVER		RELATIVE		OTHER REPRESENTATIVE		PATIENT		
20. Treatment pack number Take lowest available number treatment pack	вох					PACK			
21. Date of randomisation	day		month		year				
22. Time of randomisation (24-hour clock)	ho	urs	min	utes					
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@Lshtm.ac.uk
- Send as a secure scanned document by email to haltit.data@Lshtm.ac.uk or upload a scanned copy at http://ctu-files.Lshtm.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

<u>Please note</u>: If your query is not urgent please use the normal contact details in the investigator's study file and wall posters