Trial number:		
UNIVERSITY ^{OF} BIRMINGHAM	BCT Birmingham Cirical Trials I	Unit



FOLLOW-UP FORM

IDENTIFYING DETAILS				
Trial No.:	Initials:			
Site Name:				
Date of follow-up visit:	/ Y Y Y Y			
TIME POINT				
Please indicate which trial visit this form relates to:				
6 months (after randomisation)	12 months (after randomisation)]		
CONFIRMATION OF CONSENT				
Does the participant wish to continue taking part in the CALIBRE Trial?	If no, please complete a Change of State Yes If yes, please continue to GP Visits.	us Form.		
GP VISITS				
Since the last CALIBRE Trial visit, has the participant visited their GP in relation to liver cirrhosis/ oesophageal varices? No Yes If no, please continue to Blood Tests. If no, please continue to Blood Tests. If yes, please complete the following question.				
If yes, how many times did the participant visit their GP? Please include the total for all GP visits that were required due to liver cirrhosis/oesophageal varices, since the last CALIBRE Trial visit:				
BLOOD TESTS				
Please enter results from the most recent standard care	e blood tests:			
Test	Date taken	Not available		
Sodium mmol/L	DD/MMM/YYYY			
INR (If unavailable then both PTT for patient and control needs completing):	DD/MMM/YYYY			
PTT (Patient): Seconds	DD/MMM/YYYY			

BLOOD TESTS			
Please enter results from the mo	st recent standard	care blood tests:	
PTT (Control):	. Seconds	DD/MMM/YYY	
Albumin:	g/L	DD/MMM/YYY	
Creatinine:	μmol/L	DD/MMM/YYY	
Bilirubin:	μmol/L	DD/MMM/YYY	
Alkaline Phosphatase	U/L	DD/MMM/YYYY	
eGFR:	mL/min	DD/MMM/YYYY	
Platelets:	U/L	DD/MMM/YYYY	
CONCOMITANT MEDICATIONS	a fallowing modicat	ions currently?	
Is the participant taking any of the Class	Yes	Agent	
Anticoagulants			
Anti-platelets			
ACE inhibitors			
Angiotensin II receptor blockers			
Calcium channel blockers			
Statins			
Alpha blockers			
Beta blockers			
Nitrates			
Thiazide/ loop diuretics			
Potassium-sparing diuretics			
Sildenafil (if > 1 month)			

Trial number:		CONFIDENTIAL WHEN COMPLETE	
CONCOMITANT MEDICATION	ONS		
Is the participant taking any	of the following medicat	itions currently?	
Class	Yes	Agent	
Antiviral agent			
Ursodeoxycholic acid			
Obeticholic Acid			
Fibrate			
Other (please specify)			
I confirm that all above iter	ns have been considered	ed and only those ticked are being taken No Y	′es
MEDICAL HISTORY			
Past Medical History			
Since the last CALIBRE visit,	, has the participant beer	n diagnosed with any of the following? Please tick all that	t apply:
	Yes	Yes	
Hypertension		Peripheral vascular disease	
TIA		Stroke	
Type 1 diabetes		Type 2 diabetes	
Hypercholesterolaemia		Myocardial infarction	
Trypercholesterolaethia		Wyocardia illiarction	
Please confirm all above it	ems have been consider	red and only those ticked were present No Ye	es
Smoking Status			
		If the participant has new smoked or is an ex-smok please continue to Alcoh Consumption	ker,

months?

If the participant is a current smoker or has smoked in the last 12 months, please complete the

following question.

Smoking status: Never smoked Ex-smoker Ex-smoker (<12 months) Current smoker

On average, how many cigarettes/ cigars/ cigarillos/ has the participant smoked daily for the last 12

Trial number:					
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Alcohol Consumption	
Has the participant consumed alcohol in the last 12 months?	If no, please continue to Coffee Consumption. If yes, please complete the following questions.
On average, how many units of alcohol has the participant consmonths?	umed per week over the last 12
Has an AUDIT – C assessment been undertaken? No Yes	If no or not available, please continue to Coffee Consumption. If yes, please complete the AUDIT questions below.
	mption using the AUDIT-C, and if required, AUDIT 's Audit C score is ≥ 5 , using the guidance below please paining AUDIT questions and enter the total score.
Guidance: AUDIT –C score:	

Questions	Scoring system			Scoring system			
Questions	0	1	2	3	4		
How often do you have a drink containing alcohol?	Never	Monthly or less	2-4 times per month	2-3 times per week	4+ times per week		
How many units of alcohol do you drink on a typical day when you are drinking?	1-2	3-4	5-6	7-9	10+		
How often have you had 6 or more units if female, or 8 or more if male, on a single occasion in the last year?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily		

Total AUDIT score:

Questions	Scoring system					Participant score
•	0	1	2	3	4	
How often during the last year, have you found that you were not able to stop drinking once you had started?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily	
How often during the last year, have you failed to do what was normally expected from you because of your drinking?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily	
How often during the last year, have you needed an alcoholic drink in the morning to get yourself going after a heavy drinking session?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily	
How often during the last year, have you had a feeling of guilt or remorse after drinking?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily	
How often during the last year, have you been unable to remember what happened the night before because you had been drinking?	Never	Less than monthly	Monthly	Weekly	Daily or almost daily	
Have you or somebody else been injured as a result of your drinking?	No		Yes, but not in the last year		Yes, during the last year	
Has a relative or friend, doctor or other health worker been concerned about your drinking or suggested that you cut down?	No		Yes, but not in the last year		Yes, during the last year	

Trial number:	CONFIDENTIAL WHEN COMPLETE
Coffee Consumption	
Does the participant drink coffee daily? No Yes	If no, please continue to Ultrasound Scan Report. If yes, please complete the following question.
On average, how many cups of coffee does the participant dr	ink per day?
ULTRASOUND SCAN REPORT	
Has the participant undergone an abdominal ultrasound since the last CALIBRE Trial visit?	If no, please continue to FibroScan Report. Yes If yes, please complete the following questions.
Date ultrasound was undertaken:	/ Y Y Y Y
Hepatomegaly: No Yes Not available	
Splenomegaly: No Yes If yes, size:	cm
Was the hepatic vein patent? No Yes	Not available
Was the portal vein patent? No Yes	Not available
FIBROSCAN REPORT	
Has the participant undergone a FibroScan since the last	If no, please continue to Vital Signs.
CALIBRE Trial visit?	No Yes If yes, complete the following questions and rest of form.
Date FibroScan was undertaken: DD / MM M	/ Y Y Y
Type of probe: Medium Extra Large	Both Not available
Liver stiffness measurement: kPa	Not available
Liver stiffness measurement (IQR): %	Not available
Liver stiffness measurement (valid readings):	% Not available
Controlled attenuation parameter:	dB/m Not available
Controlled attenuation parameter (IQR): dB/r	m Not available

Trial number:	CONFIDENTIAL WHEN COMPLETE
VITAL SIGNS	
Date vital signs taken: DD / MM M / Y	YYY
Blood pressure: / mmHg I	Pulse: bpm
ANTHROPOMETRIC MEASUREMENTS	
Please record measurements to the nearest whole number	r:
Date measurements taken: DD / M M / D	<u> </u>
Height: cm	Weight: kg
ECG	
	lease continue to Treatment Adherence.
Was an ECG performed? No Yes If wes n	please complete the following questions.
Was the Within normal limits Ahnormal, not clin	ically significant Abnormal, clinically significant
ECG:	
If abnormal and clinically significant, please indicate why:	
Conduction abnormalities Rhythm a	abnormalities
TREATMENT ADHERENCE	
Since the last CALIBRE Trial Visit, has participant taken CAR	RVEDILOL? No Yes
Since the last CALIBRE Trial Visit, has participant had VBL?	No Yes
If yes to carvedilol, please complete the two carvedilol of the second o	•
PARTICIPANTS WHO HAVE TAKEN CARVEDILOL	
Date first dose of carvedilol prescribed	D/MMM/YYYY
Please indicate participant percentage adherence for carved	dilol:
Always (100%)	
Almost always (75-99 %) Most (50-74%) Some (2	25-49%) Hardly any (1-24%) Never (0%)

rial number:					
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PARTICIPANTS WHO HAVE HAD VARICEAL BAND LIGATION				
Since the last CALIBRE Trial visit, has participant undergone all scheduled variceal band ligation No Yes procedures?				
If the participant has not undergone all scheduled variceal band ligation procedures, how many appointments have been missed?				
Have the participant's oesophageal varices been eradicated since the last CALIBRE Trial visit? No Yes If no , please continue to endoscopy reports question.				
If Yes , What date were the participant's oesophageal varices eradicated?				
Have the participant's oesophageal varices recurred since the last CALIBRE Trial visit?				
No Yes Unknown				
If Yes , What date did the participant's oesophageal varices reoccurred?				
DD/MMM/YYYY Unknown				
Endoscopy Reports Reports should be completed for each endoscopy undertaken since the last CALIBRE Trial visit (maximum of six), starting with the most recent				
If no, please continue to Adverse Events.				
Have endoscopies been undertaken? No Yes If yes, please complete a report for each endoscopy undertaken.				
How many endoscopies have been undertaken?				
Guidance:				
GOV2 IGV1				
GOV 1: Oesophageal varices extending to lesser curve of stomach GOV 2: Oesophageal varices IGV 1: Isolated fundal extending to gastric fundus varices IGV 2: Isolated varices in gastric antrum, body or pylorus				

Trial number:	
Endoscopy 1	

Endoscopy 1				
Date of endoscopy:	DD/MMM.	/		
		If no, ques	please continue to the oestions	sophageal varices
Banding undertaken:	No Yes Compli	cation If yes	, please complete the follo	owing question.
		=	ere was a complication con	nplete the complications
Number of bands:		durin	g procedure question	
Number of bands.		If no please continue	to the oesophageal varices	c questions
Banding undertaken:	No Yes			s questions.
Number of bands:		ij yes, pieuse compiete	e the following question.	
Please specify the num	ber of oesophageal varices pr	esent, per grade:		
Grade I:		Grade II:	Gra	de III:
Were red signs present	: No Yes]		
Were gastric varices present? No Yes If no, please continue to the next endoscopy report (if required). If yes, please complete the following questions.				
Please specify the size of	of gastric varices identified, p		complete the johowing qui	estions.
Please specify the size of	of gastric varices identified, p		Large	Not present
		er grade:		
Grade		er grade:		
Grade GOV1		er grade:		
Grade GOV1 GOV2		er grade:		
Grade GOV1 GOV2 IGV1	Small	er grade:		
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small	er grade: Medium	Large If no, please continue report.	Not present
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small	er grade: Medium	Large If no, please continue report. If yes, please comple	Not present
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small Small Small Yes Cations during the procedure	er grade: Medium	Large If no, please continue report. If yes, please comple Please tick all that approximation in the process.	Not present
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small Small Small Yes Cations during the procedure	er grade: Medium	Large If no, please continue report. If yes, please comple Please tick all that approximation in the process.	Not present

CALIBRE 6_12 FU CRF

EudraCT No.: 2018-002488-24

v3.0, 15th November 2019

Trial number:	
'ndosson 3	

Endoscopy 2						
Date of endoscopy:	Date of endoscopy: DD / MM M / Y Y Y					
		If no, ques	please continue to the oestions	sophageal varices		
Banding undertaken:	No Yes Compli	cation If yes	s, please complete the follo	owing question.		
		=	ere was a complication com	nplete the complications		
Number of bands:		aum	ng procedure question			
		If no, please continue	to the oesophageal varices	s questions.		
Banding undertaken:	No Yes		the following question.			
Number of bands:		, , , ,	, , ,			
Please specify the num	ber of oesophageal varices pr	resent, per grade:				
Grade I:		Grade II:	Gra	de III:		
Were red signs present	: No Yes]				
If no, please continue to the next endoscopy report (if required). Were gastric varices present? No Yes If yes, please complete the following questions.						
Please specify the size of	of gastric varices identified. p		, , , , , , , , , , , , , , , , , , , ,	estions.		
Please specify the size of Grade	of gastric varices identified, p		Large	Not present		
		er grade:				
Grade		er grade:				
Grade GOV1		er grade:				
Grade GOV1 GOV2		er grade:				
Grade GOV1 GOV2 IGV1	Small	er grade:				
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small	er grade: Medium	Large If no, please continue report.	Not present		
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small	er grade: Medium	Large	Not present		
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small Small Small Yes Cations during the procedure	er grade: Medium	Large If no, please continue report. If yes, please comple Please tick all that approximation in the process.	Not present		
Grade GOV1 GOV2 IGV1 IGV2 Were red signs prese	Small Small Small Yes Cations during the procedure	er grade: Medium	Large If no, please continue report. If yes, please comple Please tick all that approximation in the process.	Not present		

CALIBRE 6_12 FU CRF

EudraCT No.: 2018-002488-24

v3.0, 15th November 2019

Trial number:			CONFIDENTIAL WHEN	N COMPLETE
Endoscopy 3				
Date of endoscopy:	DD/MMM.	/ Y Y Y Y		
		lf no, quest	please continue to the oestions	sophageal varices
Banding undertaken:	No Yes Compli	cation	, please complete the follo	wing question.
			re was a complication com g procedure question	nplete the complications
Number of bands:				
Banding undertaken:	No Yes		to the oesophageal varices	questions.
		If yes, please complete	the following question.	
Number of bands:				
Please specify the numb	per of oesophageal varices pr	esent, per grade:		
Grade I:		Grade II:	Gra	de III:
Were red signs present:	No Yes]		
If no, please continue to the next endoscopy report (if required). Were gastric varices present? No Yes If yes, please complete the following questions.				
Please specify the size of gastric varices identified, per grade:				
Grade	Small	Medium	Large	Not present
GOV1				
GOV2				
IGV1				
IGV2				

GOV2					
IGV1					
IGV2					
Were red signs presen	nt: No Yes				
Were there any complic	cations during the procedure	?	No Yes	report.	e to the next endoscopy te the following questions. oply.
Y	'es			Yes	
Failed intubation			Failed banding		

CALIBRE 6_12 FU CRF

Bleeding

EudraCT No.: 2018-002488-24

v3.0, 15th November 2019

No

specify)

I confirm that all above items have been considered and only those ticked were present

Other (please

Trial number:			CONFIDENTIAL WHEN	N COMPLETE	
Endoscopy 4					
Date of endoscopy:	DD/MMM.	/ Y Y Y Y			
		If no, ques	please continue to the oestions	sophageal varices	
Banding undertaken:	No Yes Compli	cation If yes	, please complete the follo	wing question.	
			re was a complication com g procedure question	nplete the complications	
Number of bands:					
Banding undertaken:	If no, please continue to the oesophageal varices questions. Banding undertaken: No Yes If yes, please complete the following question.				
Number of bands:					
Please specify the numb	per of oesophageal varices pr	esent, per grade:			
Grade I:		Grade II:	Gra	de III:	
Were red signs present:	No Yes]			
Were gastric varices present? No Yes If no, please continue to the next endoscopy report (if required). If yes, please complete the following questions.					
Please specify the size of gastric varices identified, per grade:					
Grade	Small	Medium	Large	Not present	
GOV1					
GOV2					
IGV1					
IGV2					

CALIBRE 6_12 FU CRF EudraCT No.: 2018-002488-24

Bleeding

v3.0, 15th November 2019

No

specify)

I confirm that all above items have been considered and only those ticked were present

Other (please

Trial number:			CONFIDENTIAL WHEN	N COMPLETE
Endoscopy 5				
Date of endoscopy:	DD/MMM,	/		
		lf no, quest	please continue to the oestions	sophageal varices
Banding undertaken:	No Yes Compli	cation If yes	, please complete the follo	wing question.
			re was a complication com g procedure question	nplete the complications
Number of bands:				
Banding undertaken:	If no, please continue to the oesophageal varices questions. Banding undertaken: No Yes If yes, please complete the following question.			
Number of bands:				
Please specify the numb	per of oesophageal varices pr	esent, per grade:		
Grade I:		Grade II:	Gra	de III:
Were red signs present:	No Yes]		
Were gastric varices present? No Yes If no, please continue to the next endoscopy report (if required). If yes, please complete the following questions.				
Grade	Small	Medium	Large	Not present
GOV1				
GOV2				
IGV1				
IGV2				

GOV2				
IGV1				
IGV2				
Were red signs presen	nt: No Yes			
Were there any complic	cations during the procedure	? No Yes [report.	e to the next endoscopy ete the following questions. pply.
Υ	'es		Yes	
Failed intubation		Failed banding		

CALIBRE 6_12 FU CRF

Bleeding

EudraCT No.: 2018-002488-24

v3.0, 15th November 2019

No

specify)

I confirm that all above items have been considered and only those ticked were present

Other (please

Trial number: CONFIDENTIAL WHEN COMPLETE					
Endoscopy 6					
Date of endoscopy:	DD/MMM,	/ Y Y Y Y			
		If no, ques	please continue to the oestions	sophageal varices	
Banding undertaken:	No Yes Compli	cation If yes	, please complete the follo	wing question.	
			ere was a complication com ng procedure question	plete the complications	
Number of bands:					
Banding undertaken:	If no, please continue to the oesophageal varices questions. Banding undertaken: No Yes If yes, please complete the following question.				
Number of bands:					
Please specify the numb	per of oesophageal varices pr	esent, per grade:			
Grade I:		Grade II:	Gra	de III:	
Were red signs present:	No Yes]			
Were gastric varices present? No Yes If no, please continue to the next endoscopy report (if required). If yes, please complete the following questions.					
Please specify the size of gastric varices identified, per grade:					
Grade	Small	Medium	Large	Not present	
GOV1					
GOV2					
IGV1					

Grade	Siliali	Wicaiaiii	Laige	Not present		
GOV1						
GOV2						
IGV1						
IGV2						
Were red signs preser	Were red signs present: No Yes					
Were there any complic	ations during the procedure	? No Yes [report.	e to the next endoscopy		
			If yes, please comple Please tick all that a	te the following questions. oply.		
Y	'es		Yes			

CALIBRE 6_12 FU CRF

Failed intubation

Bleeding

EudraCT No.: 2018-002488-24

v3.0, 15th November 2019

No

Failed banding

Other (please

specify)

I confirm that all above items have been considered and only those ticked were present

Trial number:		CONFIDENTIAL WE	HEN COMPLETE	
ADVERSE EVENTS				
defined as 'expected' (i.e. are rec	ognised complication ocol? If yes, please c	ny serious adverse events that are not ns/ consequences of cirrhosis and omplete a CALIBRE SAE FORM and	No Yes	
PARTICIPANTS WHO HAVE TAKE	N CARVEDILOL			
Since the last CALIBRE Trial visit,	has the participant	experienced/ been diagnosed with: P	lease tick all that apply	
	Yes		Yes	
Gastrointestinal upset		Blurred vision		
Dizziness		Lethargy		
Rash		Headache		
Swelling in the feet or hands		Upper respiratory tract infection		
Shortness of breath		Sexual dysfunction		
Other (please specify)				
I confirm that all above items ha	ve been considered	and only those ticked were present	No Yes	
Has the participant been put onto either a split or reduced dose of carvedilol because of adverse events? No Yes If no, please continue to the question regarding stopping carvedilol. If yes, please complete the following questions.				
			Tick if not applicable	
Date carvedilol dose was split:	DD/M	MM/YYYY		
Date carvedilol was reduced:	DD/M	MM/YYYY		
Please indicate which adverse ev	ents led to the dose	of carvedilol being split or reduced: <i>Ple</i>	ase tick all that apply	
	Yes		Yes	
Gastrointestinal upset		Blurred vision		
Dizziness		Lethargy		
Rash		Headache		
Swelling in the feet or hands		Upper respiratory tract infection		
Shortness of breath		Sexual dysfunction		
Other (please specify)				

Yes

I confirm that all above items have been considered and only those ticked were present No

Trial number:	COI	NFIDENTIAL WHEN COMPLETE		
Has the participant stopped taking carvedilol because of adverse events?	No Yes	If no, please continue to the question regarding alternative treatment. If yes, please complete the following questions.		
Date carvedilol was stopped:	/ Y Y Y	Υ		
Please indicate which adverse events have led to carvedilo	ol being stopped: P	lease tick all that apply		
Yes		Yes		
Gastrointestinal upset	Blurred vision			
Dizziness	Lethargy			
Rash	Headache			
Swelling in the feet or hands	Upper respirator	y tract infection		
Shortness of breath	Sexual dysfunction	on 🔲		
Other (please specify)				
I confirm that all above items have been considered and	only those ticked	were present No Yes		
Has an alternative treatment been commenced? No	Yes	no, please continue to All Participants. yes, please complete following question.		
Which alternative treatment has been commenced?				
Propanolol Nadolol		Variceal band ligation		
Other (please specify)				
		Tick if not available		
Date alternative treatment commenced: D D / N	1 M M / Y	YYY		

PARTICIPANTS WHO	HAVE HAD VA	RICEAL BAND LIGATION	N			
Since the last CALIBRE	Trial visit, has	the participant experie	enced/ been di	agnosed with: Please tick all that apply		
	Yes		Yes			
Abdominal pain		Banding-related dysphagia		Other (please specify):		
Has treatment with vastopped because of ac	dverse events?	INO I	Yes	If no, please continue to All Participants. If yes, please complete the following questions.		
Please indicate which	adverse event	s have led to variceal b	and ligation be	ing stopped:		
	Yes		Yes			
Abdominal pain		Banding-related dysphagia		Other (please specify):		
I confirm that all above	vo itoms have	been considered and o	nly those ticks	ed were present No Yes		
T COMMINICAL AN ADOV	ve items mave	been considered and o	illy those ticks			
Has an alternative tre	If no, please continue to All Participants. Has an alternative treatment been commenced? No Yes If yes, please complete the following question.					
Which alternative trea	atment has be	en commenced?				
Carvedilol		Propranolol [Nadolol		
Other (please specify)						
Tick if not available						
Date alternative treatment commenced: DD / MM M / YYYY						

Trial number:					
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ALL PARTICIPANTS					
Variceal Bleeding					
Guidance: The first variceal bleed is defined as hematem haemorrhage and at least a 2 g/L reduction in The definition includes bleeding from banding	haemoglobin within 2				
Has the participant experienced varic the last CALIBRE Trial visit	eal bleeding since	No 🗌	Yes	If no, please continue to As	
Was the bleeding banding related		No 🗌	Yes		
How many episodes of variceal bleeding has the participant experienced	Please complete t	he dates be	elow with m	nost recent first	Not available
Episode 1	D D	/ M M	M/	Y Y Y Y	
Episode 2	DD/MMM/YYYY				
Episode 3	D D	/ M M	M/	Y Y Y Y	
Episode 4	D D	/ M M	M/	YYYY	
Since the last CALIBRE Trial visit, has the participant been hospitalised due to variceal No Yes If yes, please complete the following questions.					
Since the last CALIBRE Trial visit, how many times has the participant been hospitalised due to variceal bleeding?		hospitalise Please includ	d? de the total j ed due to val	al, has the participant been for all hospitalisations that riceal bleeding, since the last	
Since the last CALIBRE Trial visit, was the participant admitted to intensive care due to variceal bleeding?	No Yes	intensive ca Please includ	are? de the total j	did the participant stay in for all intensive care days that the bleeding, since the last CALIBRA	
Since the last CALIBRE Trial visit, did the participant require additional outpatient visits due to variceal bleeding?	No Yes	participant Please includ	require? de the total j	tional outpatient visits did the for all outpatient visits that were bleeding, since the last CALIBRI	re

Trial number:	CONFIDENTIAL WHEN COI	MPLETE
Ascites:		
Grade 2: Moderate as	ading system detectable only by ultrasound scites manifested by moderate symmetrical distension of the abdomen ss ascites with marked abdominal distension	
Has the participant experi ascites since the last CALI		
How many episodes of new or worsening ascites has the participant experienced	Please complete the dates below with most recent first, grade and if new or worsening	Not available
Episode 1	DD/MMM/YYY	
Grade	Grade 1: Grade 2: Grade 3:	
New or Worsening	New: Worsening:	
Episode 2	DD/MMM/YYY	
Grade	Grade 1: Grade 2: Grade 3:	
New or Worsening	New: Worsening:	
Episode 3	DD/MMM/YYY	
Grade	Grade 1: Grade 2: Grade 3:	
New or Worsening	New: Worsening:	
Episode 4	DD/MMM/YYYY	
Grade	Grade 1: Grade 2: Grade 3:	
New or Worsening	New: Worsening:	

Since the last CALIBRE Trial visit, how many times has the participant been hospitalised due to Ascites?

been hospitalised due to Ascites?

Since the last CALIBRE Trial visit, has the participant

ant No Yes

If no, please continue to spontaneous bacterial peritonitis.

If yes, please complete the following questions.

How many days in total, has the participant been hospitalised?

Please include the total for all hospitalisations that have occurred due to Ascites, since the last CALIBRE Trial visit:

	•

Trial number:		CONFIDENTIAL WHEN COMPLETE	
Since the last CALIBRE Trial visit, was the participant admitted to intensive care due to Ascites?	No Yes	If yes, how many days did the participant stay in intensive care? Please include the total for all intensive care days that were required due to Ascites, since the last CALIBRE Trial visit:	
Since the last CALIBRE Trial visit, did the participant require additional outpatient visits Ascites?	No Yes	If yes, how many additional outpatient visits did the participant require? Please include the total for all outpatient visits that were required due to Ascites, since the last CALIBRE Trial visit:	

Trial number:		CONFIDENTIAL WHEN	COMPLETE	
Spontaneous Bacterial Peritonit	is:			
Since the last CALIBRE Trial visit, participant been diagnosed with bacterial peritonitis?		Yes	to Renal Dysfunction. e the following questions.	
How many episodes of spontaneous bacterial peritonitis have been diagnosed since the last CALIBRE Trial visit:	Please complete the dates be	low with most recent first	Not available	
Episode 1		MM/YYYY		
Episode 2	DD/M	MM/YYYY		
Episode 3	DD/M[MM/YYYY		
Episode 4	DD/M	MM/YYYY		
Since the last CALIBRE Trial visit, has the participant been hospitalised due to spontaneous No Yes If yes, please complete the following questions.				
Since the last CALIBRE Trial visit, h many times has the participant be hospitalised due to spontaneous bacterial peritonitis?		How many days in total, has the particip hospitalised? Please include the total for all hospitalis have occurred due to spontaneous bacto peritonitis, since the last CALIBRE Trial v	ations that	
Since the last CALIBRE Trial visit, was the participant admitted to intensive care due to	No Yes	If yes, how many days did the participar intensive care? Please include the total for all intensive were required due to spontaneous bacte peritonitis, since the last CALIBRE Trial v	care days that erial	

CALIBRE 6_12 FU CRF

spontaneous bacterial

Since the last CALIBRE Trial visit,

additional outpatient visits due

did the participant require

to spontaneous bacterial

peritonitis?

peritonitis?

EudraCT No.: 2018-002488-24

If yes, how many additional outpatient visits did the

Please include the total for all outpatient visits that

were required due to spontaneous bacterial

peritonitis, since the last CALIBRE Trial visit:

participant require?

Yes

Trial number:		CONFIDENTIAL V	VHEN COMPLETE		
Renal Dysfunction:					
Since the last CALIBRE Trial visit, has the Hepatorenal Syndrome per Internation		NOL I YEST I	If no, please cont Hepatocellular Co If yes, please com following questio	arcinoma. aplete the	
How many episodes of Hepatorenal Syndrome have been diagnosed since the last CALIBRE Trial visit:	Please complete the da	tes below with most recent first		Not available	
Episode 1	D)/MMM/YYYY			
Episode 2	DD)/MMM/YYYY			
Episode 3	D	/MMM/YYYY			
Episode 4	D)/MMM/YYYY			
Since the last CALIBRE Trial visit, has the hospitalised due to Hepatorenal Syndron	NOT	Yes	tinue to Hepatocellu		
How many days in total, has the participant been hospitalised? Hepatorenal Syndrome? How many days in total, has the participant been hospitalised? Please include the total for all hospitalisations that have occurred due to Hepatorenal Syndrome, since the last CALIBRE Trial visit:					
Since the last CALIBRE Trial visit, was the participant admitted to intensive care duto Hepatorenal Syndrome?		If yes, how many days did the partici intensive care? Please include the total for all inten- were required due to Hepatorenal S last CALIBRE Trial visit:	sive care days that		
Since the last CALIBRE Trial visit, did the participant require additional outpatient visits due to Hepatorenal Syndrome?	: No Yes	If yes, how many additional outpati participant require? Please include the total for all outpo were required due to Hepatorenal S last CALIBRE Trial visit:	atient visits that		
Guidance: Diagnostic criteria of hepatorenal syndrome (HRS) HRS-AKI	type of acute kidney injury (AKI)	in patients with cirrhosis			

- Diagnosis of cirrhosis and ascites
- Diagnosis of AKI according to ICA-AKI criteria*
- No response after 2 consecutive days of diuretic withdrawal and plasma volume expansion with
 - albumin 1 g/kg bodyweight
- Absence of shock
- No current or recent use of nephrotoxic drugs (NSAIDs, aminoglycosides, iodinated contrast media, etc)
- No macroscopic signs of structural kidney injury#, defined as:
 - o absence of proteinuria (>500 mg/day)
 - o absence of microhaematuria (>50 RBCs per high power field)
 - o normal findings on renal ultrasonography

*Patients who fulfil these criteria may still have structural damage such as tubular damage. Urine biomarkers will become an important element in making a more accurate differential diagnosis between HRS and acute tubular necrosis.

ICA, International Club of Ascites; NSAIDs, non-steroidal anti-inflammatory drugs; RBCs, red blood cells.

#Increase in sCr ≥0.3 mg/dL (≥26.5 µmol/L) within 48 h; or a percentage increase sCr ≥50% from baseline which is known, or presumed, to have occurred within the prior 7 days. (A value of sCr obtained in the previous 3 months, when available, can be used as baseline sCr. In patients with more than one value within the previous 3 months, the value closest to the admission time to the hospital should be used. In patients without a previous sCr value, the sCr on admission should be used as baseline).

CALIBRE 6_12 FU CRF EudraCT No.: 2018-002488-24 v3.0, 15th November 2019

Trial number:			
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Hepatocellular Carcinoma (HCC):				
Since the last CALIBRE Trial visit, has the partici diagnosed with HCC?	ipant been	No Yes	If no, please continue to Encephalopathy. If yes, please complete questions.	·
Date DD/MM/M/YYYY		Not available		
BCLC stage: Stage 0 Stage	Α 🗌	Stage B	Stage C	Stage D
When was the participant diagnosed?	\mathbb{M}	/ Y Y Y	Υ	
Please indicate which treatment (if any) that th	ne participar	nt is on: <i>Please tick</i>	all that apply	
Yes		Yes		Yes
Loco-regional therapy Sys	temic thera	ру	Radiation therapy	
Surgery	ner (<i>please</i> ecify)			
I confirm that all above items have been consi	dered and	only those ticked w	vere present No	Yes
Since the last CALIBRE Trial visit, has the participant been hospitalised No Yes due to HCC? If no, please continue to Hepatic Encephalopathy. If yes, please complete the following questions.				
How many days in total, has the participant been hospitalised? The participant been hospitalised due to HCC? How many days in total, has the participant been hospitalised? Please include the total for all hospitalisations that have occurred due to HCC, since the last CALIBRE Trial visit:				
Since the last CALIBRE Trial visit, was the participant admitted to intensive No care due to HCC?	Yes	intensive care? Please include the to	ays did the participant s tal for all intensive care do HCC, since the last CALIBI	ays that
Since the last CALIBRE Trial visit, has the participant required additional No outpatient visits due to HCC?	Yes	the participant req	dditional outpatient vis uire? tal for all outpatient visits HCC, since the last CALIBI	that

Guidance:

BCLC Staging System

- Stage 0: The tumour is less than 2 cm, the person feels well (performance status 0) and the liver is working normally
- Stage A: There is a single tumour less than 5 cm, or up to 3 tumours all less than 3cm. The person feels well and is active (performance status 0) and the liver is working well
- Stage B: There are many tumours in the liver, but the person feels well (performance status 0) and the liver is working well
- Stage C: The cancer has spread into the blood vessels, lymph nodes or other body organs. Or the person does not feel well (performance status 1 or 2). The liver is still working.
- Stage D: There is severe liver damage or the person is not well and needs help in being looked after (performance status 3 or 4)

Trial number:				

Hepatic Encephalopathy:							
Since the last CALIBRE Trial participant been diagnosed encephalopathy?	<u> </u>						
How many episodes of hepatic encephalopathy have been diagnosed since the last CALIBRE Trial visit:	Please complete the dates below with most recent first, grade and if new or worsening	Not available					
Episode 1	DD/MMM/YYYY						
Grade	Grade 1: Grade 2: Grade 3: Grade 4:						
New or Worsening	New: Worsening:						
Episode 2	DD/MMM/YYYY						
Grade	Grade 1: Grade 2: Grade 3: Grade 4:						
New or Worsening	New: Worsening:						
Episode 3	DD/MMM/YYYY						
Grade	Grade 1: Grade 2: Grade 3: Grade 4:						
New or Worsening	New: Worsening:						
Episode 4	DD/MMM/YYYY						
Grade	Grade 1: Grade 2: Grade 3: Grade 4:						
New or Worsening	New: Worsening:						
I confirm that all above items have been considered and only those ticked were present No Yes							
If no, please continue to Liver Transplant question. Since the last CALIBRE Trial visit, has the participant been hospitalised due to hepatic No Yes encephalopathy? If yes, please complete the following questions.							
Since the last CALIBRE Trial visit, how many times has the participant been hospitalised due to hepatic encephalopathy? How many days in total, has the participant been hospitalised? Please include the total for all hospitalisations that have occurred due to hepatic encephalopathy, since the last CALIBRE Trial visit:							

CALIBRE 6_12 FU CRF EudraCT No.: 2018-002488-24 v3.0, 15th November 2019

Trial number:	CONFIDENTIAL WHEN COMF	PLETE			
was the participant admitted to intensive care due to hepatic	If yes, how many days did the participant stay in intensive care? Please include the total for all intensive care days that were required due to hepatic encephalopathy, since the last CALIBRE Trial visit:				
did the participant require additional outpatient visits due	If yes, how many additional outpatient visits did the participant require? Please include the total for all outpatient visits that were required due to hepatic encephalopathy, since the last CALIBRE Trial visit:				
	shortened attention span; impaired performance of additic time or place; subtle personality change; inappropriate be erbal stimuli; confusion; gross disorientation				
Liver Transplant:					
Since the last CALIBRE Trial visit, has the participant ha transplant?	questions to	se complete the			
Specify date of liver transplant below					
Date DD/MMM/YY	YY				
CRF completed by: You must have signed the CALIBRE site training log and CALIBRE delegation log	Name:(please print)				
Date: DD / MM M / V V	Y Signature:				

CALIBRE 6_12 FU CRF v3.0, 15th November 2019 EudraCT No.: 2018-002488-24