

Ambition Trial Coordinating Centre Private Bag 320, Princess Marina Hospital Gaborone, Botswana



Laboratory Work Practice Document: 3 (Gaborone) Sample Processing and Storage								
Title of studyHigh Dose AMBISOME <sup>©</sup> on a Fluconazole Backbone for Cryptococcal Meningitis Induction Therapy in sub-Saharan Africa: A Phase III Randomized Controlled Non-inferiority Trial								
Acronym	Ambition-cm – AMBIsome Therapy Induction OptimizatioN							
ISRCTN No.:	ISRCTN72509687							
WPD Current version	VPD Current version Version 1.3 22/06/2020							
Author(s)	Kwana Lechiile Lab Scientist	Heaven Feeling	22/06/2020					
	Timothée Boyer Chammard Clinical Advisor		22/06/2020					
Reviewer(s)	r(s) David Lawrence Lead Clinician 22/06/2020							
Approved by	Joseph Jarvis Cl J.N.J. 22/06/2020							

Revision History:							
Version Number	Effective Date	Reason for Change					
1.0	14/06/2018	First version					
1.1	22/08/2018	Addition of India ink storage at D14					
1.2	10/09/2019	Addition of PAXgene Specimen Collection and Storage					
1.3	22/06/2020	Addition of PBMC isolation					



#### Purpose

This document describes the day to day laboratory duties

#### References

Ambition Phase III Trial Protocol

#### Materials

Lab WPD 1 – Laboratory equipment Lab WPD 2 – Quantitative Cryptococcal cultures Lab WPD 7 – Quantitative Cryptococcal PCR sub-study Lab WPD 8 – Cryptococcal Semi-Quantitative Antigen Lab WPD 9 - PAXgene Specimen Collection and Storage Clinical WPD 15- Timing of evaluation and tests Clinical WPD 20 – Quantitative PCR Sub-study

#### Appendices

Lab Processing Chart: Gaborone

#### Section A – Lab maintenance, quality control

Monitoring of freezer and fridge temperatures, reagents/agar expiration dates, working order of autoclaves, incubators and other laboratory equipment, should all be performed in line with on-site laboratory quality control and health and safety procedures

Weekly stock check of lab consumables for trial. Inform trial team on-site and trial manager when replacements required. See Laboratory WPD 1 – Laboratory equipment



#### Section B – Samples required

All results should be recorded with patient study number, date, and study day.

All results should be communicated to the study team.

#### – Study bloods

The following must be performed according to the trial protocol

- CD4 at baseline if not done within 3 months
- FBC and ALT on days 1, 7, 14, 28
- Creatinine, electrolytes on days 1, 3, 5, 7, 10, 12, 14, 28
- Biosynex CryptoPS CrAg on whole blood on D1

(please see Lab WPD 8 – Cryptococcal Semi-Quantitative Antigen)

- qPCR samples (4ml of blood EDTA tubes): D1 (pre-dose), 24 hours post-dose, D3, D7 and D14, (please see Appendix, Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study)
- PAXgene tube on D1 (2.5mL of blood) (please see Appendix and Lab WPD 9 - PAXgene Specimen Collection and Storage)
- PBMC isolation on D1 (20mL of blood)

#### – CSF

- India Ink examination and Cryptococcal antigen (IMMY CrAg) on D1
- Biosynex CryptoPS CrAg on CSF on D1

(please see Lab WPD 8 – Cryptococcal Semi-Quantitative Antigen)

- CSF cell count and differential, protein, glucose and routine culture on D1 for all patients and additional LPs, as indicated
- Quantitative fungal culture will be conducted on D1, 7, and 14
- CSF for qPCR (1ml): D1 (pre-dose), D7 and D14 (please see Appendix, Lab WPD7 & Clinical WPD 20 - Quantitative PCR Sub-study)
- PAXgene tube on D1 (2.5mL of CSF) (please see Appendix and Lab WPD 9 - PAXgene Specimen Collection and Storage)



Event Schedule	Screening	ening Week 1					Week 2						Wk 4			
Study Day	≤D0	D1	D2	D3	D4	D5	D6	D7	D8	D9	D10	D11	D12	D13	D14	
Clinical labs																
HIV testing (if status unknown)*		х														
Pregnancy Test (Urine/Serum) (1)		х														
Full Blood Count		х						Х							х	х
CD4 count & Viral load (if needed)*		х														
ALT		х						Х							Х	х
Urea, creatinine and electrolytes		х		Х		х		Х			х		Х		Х	х
Blood for drug levels (2)		х							Х							
CSF																
Opening pressure		х						х							Х	
Cell count and differential*		х														
Protein, glucose*		х														
Routine culture *		х														
India ink examination* (3)		х														
Cryptococcal antigen* (3)		х														
Quantitative fungal culture		х						Х							Х	
CSF Drug levels (2)		Х						Х							Х	
Immune parameters (2)		X						x							x	

#### Table 1: Timings of study bloods and CSF analyses

\*Part of routine care. 1. For women of childbearing age. 2. Sub-studies at limited sites 3. India ink or CrAgrequired for inclusion.

#### Section C - Procedures for sample processing and saving - Please see Appendix: Lab Processing Chart

AMBITION-*cm* 

ne Therapy Induction OptimizatioN

#### <u>Day 1 (or 0)</u>

#### Required samples:

- 2 yellow (or red) top (1 for biochemistry, 1 for serum storage)
- ≥3 purple top (1 for FBC, 1 for whole blood storage, 1 for plasma and buffy coat storage and other samples may be for CD4, viral load and resistance testing)
- CSF sample in 3 x white top tube (5 ml each)
- qPCR samples
- (Please see WPD 20 Quantitative PCR sub-study)
- 2 PAXgene tubes (1 for blood, 1 for CSF)

(Please see WPD 9 - PAXgene Specimen Collection and Storage)

- 20mL lithium heparin blood for PBMC isolation

#### DAY 1 CSF SAMPLES

NB CSF must be processed within 4 HOURS of receipt

- 1. Perform protein and glucose on CSF
- 2. Perform India Ink, CrAg (if required), cell count (on day 0/1 sample)
- 3. Plate out day 1 unspun CSF dilutions 0-4 (neat, one in 10, 100, 1,000 and 10,000) for quantitative culture (see WPD 2)

Store isolate once grown in 2 Microbank vials.

- 4. Vortex CSF and save: divide into 3 x 2ml Cryovials min 200  $\mu$ L each. Label and place into freezer box.
- 5. qPCR SAMPLES: Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study
- 6. Biosynex CryptoPS CrAg: Please see Lab WPD 8 Cryptococcal semi-quantitative antigen
- 7. PAXgene tube: Please see WPD 9 PAXgene Specimen Collection and Storage

TB culture only if clinically indicated (will be performed in TB lab, and is not mandated by trial protocol)

#### DAY 1 BLOOD SAMPLES

- Take blood for CD4, viral load and/or resistance testing (if indicated), whole blood, plasma and buffy coat (all purple tops) and serum (yellow or red top). Total required = 2 yellow or red top (4 ml) and ≥3 purple tops (4 ml each)
- 2. Perform FBC, ALT, Creatinine and electrolytes on Day 1
- 3. <u>WHOLE BLOOD</u>: take 2x cryovials and pipette 900 uL of WHOLE BLOOD (unspun sample) into each (should be mixed well in purple top tubes prior to transfer). Label and place into whole blood storage box.

4. <u>PLASMA AND BUFFY COAT</u>: Spin blood from purple top tube at 800xg for 15 minutes. Avoiding the buffy coat and red cells at the bottom, remove as much of the plasma as possible using a sterile Pasteur pipette and divide into 2 cryovials.

Buffy coat is the small region between the plasma and red blood cells. After removing the plasma, use a sterile Pasteur pipette to carefully aspirate the buffy coat in a circular movement trying not to disturb the red blood cell layer then store in a single cryovial

AMBITION-cm

ne Therapy Induction OptimizatioN

- 5. <u>SERUM</u>: Spin yellow (or red) top sample at 800xg for 15 minutes. In the yellow top, gel now separates cells from serum. Use a sterile Pasteur pipette to remove as much of the serum possible and divide into 2 cryovials.
- 6. qPCR SAMPLES: Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study
- 7. Biosynex CryptoPS CrAg: Please see Lab WPD 8 Cryptococcal semi-quantitative antigen
- 8. PAXgene tube: Please see WPD 9 PAXgene Specimen Collection and Storage
- 9. PBMC Isolation: 20mL blood in Lithium Heparin tube

#### 24 hours post-dose

#### Required samples:

- <u>aPCR blood: Please see WPD 20 – Quantitative PCR Sub-study</u>

#### <u>Day 3</u>

#### **Required samples:**

- 1 purple top (for plasma and buffy coat)
- 1 yellow/red top tube (4ml)
- qPCR samples

(Please see Lab WPD7 & Clinical WPD 20 – Quantitative PCR sub-study)

#### DAY 3 BLOOD SAMPLES

- 1. Perform urea, creatinine and electrolytes on Day 3
- 2. PLASMA AND BUFFY COAT
- 3. qPCR SAMPLES: Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study

#### <u>Day 5</u>

#### **Required samples:**

- 1 yellow/red top tube (4ml)

#### DAY 5 BLOOD SAMPLES

1. Perform urea, creatinine and electrolytes on Day 5

#### <u>Day 7</u>

#### Required samples:

- 2 purple top (1 for FBC, 1 for plasma and buffy coat storage)
- 1 yellow/red top tube (4ml)
- CSF sample in 2 x white top tubes (5 ml each)
- qPCR samples (Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR sub-study)

#### DAY 7 CSF SAMPLES

- 1. Perform quantitative cultures on Day 7 LP.
- 2. Save remaining CSF, divide into 3 x 2ml Cryovials.
- 3. Save Cryptococcus isolate in 2 Microbank vials if growth from Day 7 LP
- 4. qPCR SAMPLES: Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study

#### DAY 7 BLOOD SAMPLES

- 1. Perform FBC, ALT, urea, creatinine and electrolytes on Day 7
- 2. PLASMA AND BUFFY COAT
- 3. qPCR SAMPLES: Please see Lab WPD7 & Clinical WPD 20 Quantitative PCR Sub-study

#### <u>Day 10</u>

#### Required samples:

- 1 yellow/red top tube (4ml)

#### DAY 10 BLOOD SAMPLES

1. Perform urea, creatinine and electrolytes on Day 10

#### <u>Day 12</u>

#### **Required samples:**

- 1 yellow/red top tube (4ml)

#### DAY 12 BLOOD SAMPLES

1. Perform urea, creatinine and electrolytes on Day 12

#### <u>Day 14</u>

#### Required samples:

- 2 purple top (1 for FBC and 1 for plasma and buffy coat)
- 1 yellow/red top tube (4ml)
- CSF sample in 2 x white top tube (5 mls each)
- qPCR samples (including India ink storage)

(Please see Lab WPD7 & Clinical WPD 20 – Quantitative PCR sub-study)

#### DAY 14 CSF SAMPLES

- 1. Perform quantitative cultures on Day 14 LP. Save remaining CSF, divide into 3 x 2ml Cryovials.
- 2. Save Cryptococcus isolate in 2 Microbank vial if growth from Day 14 LP
- 3. qPCR SAMPLES: Please see WPD 20 Quantitative PCR Sub-study

#### DAY 14 BLOOD SAMPLES

- 1. Perform FBC, ALT, urea, creatinine and electrolytes on Day 14
- 2. PLASMA AND BUFFY COAT
- 3. qPCR SAMPLES: Please see WPD 20 Quantitative PCR Sub-study

#### <u>Day 28</u>

#### Required samples:

- 1 purple top (for FBC)
- 1 yellow/red top tube (4ml)

#### **DAY 28 BLOOD SAMPLES**

1. Perform FBC, ALT, urea, creatinine and electrolytes on Day 28



#### PROCESSING AND LABELLING

**SPINNING:** For all samples use 800xg for 15 mins, unless stated otherwise on LPC.

**LABELLING**: Each sample must be labelled with study number (e.g. AMB 1-1-001), nature of sample (e.g. CSF/ plasma etc), study day (e.g. d3), date and time (time specimen taken, <u>not</u> time specimen received in lab).

The study day is derived from **the first day the patient receives study drugs**. That is day 1. That is not necessarily the day of hospital admission.

**STORAGE:** Cryovials containing samples should be stored in appropriately labelled cryoboxes. Cryoboxes should be stored in -80°C freezer

**CRYPTOCOCCAL ISOLATE STORAGE:** Cryptococcal isolates are stored on Microbank beads in an appropriately labelled cryobox and stored in -80°C freezer.

#### PROCEDURE FOR STORING ON MICROBANK BEADS:

- 1. Using a permanent marker, label with the study number, date and day of study (e.g. AMB 1-1-001, 12.10.15, d7)
- 2. Under aseptic conditions open the screw cap cryovial.
- 3. Inoculate the cryopreservative fluid with young colonial growth (18-24 hours) picked from a pure culture to approximately a 3-4 McFarland standard.
- 4. Close vial tightly and invert 4-5 times to emulsify organism. DO NOT VORTEX!
- 5. At this point the microorganisms will be bound to the porous beads. The excess cryopreservative fluid should be well aspirated leaving the inoculated beads as free of liquid as possible. Close the vial finger tight.
- 6. Record on the storage log.
- 7. Store the inoculated cryovial in -80°C freezer.



#### Section D - Summary of samples to be saved

Day 1:	Serum (stored in 2 Cryovials)
	Plasma (divided into 2 Cryovials)
	Buffy coat (stored in 1 cryovial)
	Whole blood (divided into 2 cryovials)
	CSF (divided into 3 Cryovials)
	Crypto isolate (in 2 Microbank vials)
	qPCR whole blood (stored in 1 ceramic beaded tube)
	qPCR plasma (stored in 1 eppendorf tube)
	qPCR CSF supernatant (stored in 1 cryovial)
	qPCR CSF pellet (stored in 1 cryovial)
	Blood PAXgene tube
	CSF PAXgene tube
	PBMCs
24HR:	qPCR whole blood (stored in 1 ceramic beaded tube)
	qPCR plasma (stored in 1 eppendorf tube)
Day 3:	Plasma (divided into 2 Cryovials)
	Buffy coat (stored in 1 cryovial)
	qPCR whole blood (stored in 1 ceramic beaded tube)
	qPCR plasma (stored in 1 eppendorf tube)
Day 7:	Plasma (divided into 2 Cryovials)
	Buffy coat (stored in 1 cryovial)
	CSF (divided into 3 Cryovials)
	Crypto isolate (in 2 Microbank vials – if growth from day 7 LP)
	qPCR whole blood (stored in 1 ceramic beaded tube)
	qPCR plasma (stored in 1 eppendorf tube)
	qPCR CSF supernatant (stored in 1 cryovial)
	qPCR CSF pellet (stored in 1 cryovial)
Day 14:	Plasma (divided into 2 Cryovials)
	Buffy coat (stored in 1 cryovial)
	CSF (divided in 2 Cryovials)
	Crypto isolate (in 1 Microbank vial (if growth from Day 14 LP)
	qPCR whole blood (stored in 1 ceramic beaded tube)
	qPCR plasma (stored in 1 eppendorf tube)
	qPCR CSF supernatant (stored in 1 cryovial)
	qPCR CSF pellets (stored in 2 cryovial)
On D1, D3,	D7 and D14 for qPCR or semi-quantitative CrAg sub-studies please refer to the corresponding

### Lab Processing Chart - Botswana

### AMBITION-cm

High Dose AMBISOME on a Fluconazole Backbone for Cryptococcal Meningitis Induction Therapy in sub-Saharan Africa: A Phase 3 Randomised Controlled Non-inferiority Trial

(As per Protocol version 2.1: 07/11/2017)

Protocol Version: 2.1 07/11/2017

Original LPC: Version 1.0, 14/06/2018

Revised/Updated: Version 1.1 03/07/2018: addition of India ink storage at D14 Version 1.2 10/09/2019 : addition of PAXgene specimen collection and storage at D1

Laboratory Technologist: Kwana Lechiile Em

Email: klechiile@bhp.org.bw

Phone: +267 3902671

Lead Clinician: Dr David Lawrence

Clinical Advisor: Dr Timothee Boyer-Chammard



# Laboratory Working Practice Document 3:

	TABLE 6.1. Event Schedule	Screening			V	Veek	1			Week 2						Wk 4	Wk 6	Wk 8	Wk 10	Wk 16	
Table 1: Schedule of Events	Study Day	≤D0	D1	D2	D3	D4	D5	D6	D7	D8	D9	D10	D11	D12	D13	D14					
	Consent forms																				
	PIS and signed consent	Х	Х																		
	Follow up																				
	Screening and randomisation	Х	Х																		
	Clinical review		Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х					
	Outpatient follow-up															_	Х	х	Х	Х	
	Week 16 telephone															_					х
	Clinical labs																				
	HIV testing (if status unknown)*		Х																		
	Pregnancy Test (Urine/Serum) (2)		Х																		
	Full Blood Count		Х						Х							х	Х				
	CD4 count & Viral load (if needed)*		Х																		
	ALT		Х						Х							х	Х				
	Urea, creatinine and electrolytes		Х		Х		Х		Х			Х		Х		Х	Х				
	qPCR (4)		Х						Х							Х					
	Clinical evaluation																				
	Chest X-ray (1)		Х																		
	CSF																				
	Opening pressure		Х						Х							Х					
	Cell count and differential*		Х																		
	Protein, glucose*		Х																		
	Routine culture *		Х																		
	India ink examination* (3)		Х																		
	Cryptococcal antigen* (3)		Х																		
	Quantitative fungal culture		Х						Х							Х					
	Immune parameters and qPCR(4)		Х						Х							Х					

\*Part of routine care. 1. If clinically indicated. 2. For women of childbearing age. 3. India ink or cryptococcal antigen required for inclusion. 4. As part of sub-studies at limited sites.

Page 12

AMBITION-CM

# Laboratory Working Practice Document 3:

#### Table 2: Specimen Log

Evaluation	Specimen	Tests	Special Notes
Haematology	EDTA	FBC	All patients, when applicable
CD4	EDTA	CD4 count	Should be performed if it has not been done within 3 months or if there is
			reason to suspect it is required.
Viral Load	EDTA	Viral load	Should be performed if it has not been done within 3 months or if there is
			reason to suspect it is required.
Resistance	EDTA	Resistance testing	If patient is on ART, has a detectable viral load, and reports good
			adherence or their ART history gives reason to suspect resistance, then
			resistance testing will be performed if possible at the site.
Chemistry	Yellow/Red Top Tube	ALT, Creatinine, Electrolytes	All patients, when applicable
Whole Blood	EDTA	For further analysis and sub-studies	Storage at -80°C until shipment
Plasma	EDTA	For further analysis and sub-studies	Storage at -80°C until shipment
Serum	Yellow/Red Top Tube	For further analysis and sub-studies	Storage at -80°C until shipment
Whole blood for	EDTA	For qPCR sub-study in blood	Storage at -80°C until shipment
qPCR			
Chemistry – CSF	CSF tube	Protein, Glucose	All patients, when applicable
Microbiology –	CSF tube	India Ink, CrAg LFA, Gram stain, Cell count and differentiation,	Store isolate and any unused sample at -80°C until shipment
CSF		Routine culture (for D1 CSF and if clinically indicated thereafter)	
		Quantitative cryptococcal culture (for all CSF)	
CSF for qPCR	CSF tube	For qPCR sub-study in CSF	Storage at -80°C until shipment
Blood PAXgene	PAXgene tube	For PAXgene tube sub-study	Storage immediately at 18-25°C, then at -20°C or at -80°C
CSF PAXgene	PAXgene tube	For PAXgene tube sub-study	Storage immediately at 18-25°C, then at -20°C or at -80°C
PBMCs	Lithium heparin	For PBMC isolation	As per local SOP.

Page 13

#### **Classification of samples:**

**Routine Tube:** Part of routine clinical care, tests performed by usual laboratory and not required to be in an ISO accredited lab. Not curated by Ambition.

Monitoring Tube: As part of protocol driven monitoring, stored if samples remain and curated by Ambition.

**Study Tube:** For specified sub-studies or storage for future research. Curated by Ambition.

AMBITION-CM

### Laboratory Working Practice Document 3:

### Sample Processing and Storage

#### Table 3: Study lay-out

Visit	Specimen	Tests Conducted	Storage	Special Notes
		A number of samples on Day 1 are routine test	ts which may be handled by other laborator	ies
	1	*Store any unused samples* *Aim f	for a <u>minimum</u> of 5ml CSF from each LP*	1
Day 1	CSF (5ml) x 3	Routine Tube 1: perform protein and glucose, cell count and differential, CrAg LFA, India Ink, Gram stain, routine culture	No storage for study required.	May need to separate into two: biochemistry and microbiology
		Study Tube 1: perform quantitative cryptococcal culture in duplicates	Vortex and divide remaining CSF into 3x2ml cryovials (min. 200µl each) and store at -80°C	Store isolate once grown in 2 Microbank vials and keep at -80°C
		Study Tube 2: Sample for storage	General CSF storage: Unspun sample	Store at -80°C
			qPCR sub-study: centrifuge CSF at 2000g for 10 min and store supernatant and pellet in different cryovials at -80°C	
	CSF PAXgene tube	Immediately after adding CSF, gently invert the PAXgene tube 8-10 times.	Immediately after collecting specimens, store the PAXgene tube at 18-25°C for a minimum of 2 hours and a maximum of	When storing in the freezer the preference is to store on a wire rack: do not freeze tubes upright in a
		Samples do not require any processing.	72 hours. The PAXgene tubes can then be stored at -20°C or below. If you would prefer to store at -80°C then it will be necessary to first store at - 20°C for at least 24 hours before transferring to the deep freeze.	Styrofoam tray as this may cause the tubes to crack. If you cannot store on a wire rack then keep the tubes in a freezer bag and store them in a safe, secure position in the freezer.
	EDTA (4ml)	Optional Routine Tube 1 and 2: CD4 and/or VL/Resistance Test (one tube for each test)		
		Monitoring Tube 1: FBC	After FBC, spin at 800 <sup>×</sup> g for 15min and store remaining plasma as below	If being performed in research laboratory.

		Study Tube 1: Whole blood	Invert 8-10 times and pipette into 2x2ml cryovials (min. 900µl each) and store at - 80°C	
		Study Tube 2: Spin tube at 800 <sup>x</sup> g for 15min and	Divide plasma into 2x2ml cryovials.	Store at -80°C
		remove plasma	Remove buffy coat and put into 1 x 2ml cryovial	
		Study Tube 3: qPCR sub-study	Remove 1.2ml and store in 2ml cryovial	
			remaining 3.8ml at 2000g for 10min and store plasma in 1.5ml Eppendorf tube	
	Yellow/Red Top Tube (4ml) x 2	Monitoring Tube 1: ALT, urea, creatinine, and electrolytes	Spin and store remaining serum as below.	If being performed in research laboratory.
		Study Tube 1: Spin tube at 800 <sup>x</sup> g for 15min and remove serum	Divide serum into 2x2ml cryovials and store at -80°C	
	Blood PAXgene tube	Immediately after adding the blood, gently invert the PAXgene tube 8-10 times.	Immediately after collecting specimens, store the PAXgene tube at 18-25°C for a minimum of 2 hours and a maximum of	When storing in the freezer the preference is to store on a wire rack: do not freeze tubes upright in a
		Samples do not require any processing.	72 hours. The PAXgene tubes can then be stored at -20°C or below.	Styrofoam tray as this may cause the tubes to crack. If you cannot store on a wire rack then
			If you would prefer to store at -80°C	keep the tubes in a freezer bag and store
			then it will be necessary to first store at - 20°C for at least 24 hours before	them in a safe, secure position in the freezer.
			transferring to the deep freeze.	
	Lithium heparin (20ml)	PBMC Isolation	As per local SOP	
24 hours	EDTA (4ml) x 1	Study Tube 1: qPCR sub-study	Remove 1.2ml and store in 2ml cryovial	qPCR tube is collected within 1 h of t23
post-dose			remaining 3.8ml at 2000g for 10min and	
			store plasma in 1.5ml Eppendorf tube	
Day 3	Yellow/Red Top	Monitoring Tube 1: Urea, creatinine and electrolytes	Spin and store remaining serum.	If being performed in research
	Tube (4ml)			laboratory.

	EDTA (4ml) x 2	Study Tube 1: Spin tube at 800 <sup>x</sup> g for 15min and remove plasma	Divide plasma into 2x2ml cryovials. Remove buffy coat and put into 1 x 2ml cryovial	Store at -80°C
		Study Tube 2: qPCR sub-study	Remove 1.2ml and store in 2ml cryovial containing ceramic beads; centrifuge remaining 3.8ml at 2000g for 10min and store plasma in 1.5ml Eppendorf tube	
Day 5	Yellow/Red Top Tube (4ml)	Monitoring Tube 1: Urea, creatinine and electrolytes	Spin and store remaining serum.	If being performed in research laboratory.
Day 7	CSF (5ml) x 2	Study Tube 1: Perform quantitative cryptococcal culture in duplicates	Vortex and divide remaining CSF into 3x2ml cryovials (min. 200µl each) and store at -80°C	Store isolate once grown in 2 Microbank vials and keep at -80°C
		Study Tube 2: Sample for storage	General CSF storage: Unspun sample qPCR sub-study: centrifuge CSF at 2000g for 10min and store supernatant and pellet in different cryovials at -80°C	Store at -80°C
	EDTA (4ml) x 3	Monitoring Tube 1: FBC	After FBC, spin at 800 <sup>x</sup> g for 15min and store remaining plasma as below	If being performed in research laboratory.
		Study Tube 1: Spin tube at 3500rpm/ 800 <sup>x</sup> g for 15min and remove plasma	Divide plasma into 2x2ml cryovials. Remove buffy coat and put into 1 x 2ml cryovial	Store at -80°C
		Study Tube 2: qPCR sub-study	Remove 1.2ml and store in 2ml cryovial containing ceramic beads; centrifuge remaining 3.8ml at 2000g for 10min and store plasma in 1.5ml Eppendorf tube	
	Yellow/Red Top Tube (4ml)	Monitoring Tube 1: ALT, urea, creatinine, and electrolytes	Spin and store remaining serum.	If being performed in research laboratory.
Day 10	Yellow/Red Top Tube (4ml)	Urea, creatinine and electrolytes	Spin and store remaining serum.	If being performed in research laboratory.

# Laboratory Working Practice Document 3:

Day 12	Yellow/Red Top Tube (4ml)	Monitoring Tube 1: Urea, creatinine and electrolytes	Spin and store remaining serum.	If being performed in research laboratory.
Day 14	CSF (5ml) x 2	Study Tube 1: Perform quantitative cryptococcal culture in duplicates	Vortex and divide remaining CSF into 3x2ml cryovials (min. 200µl each) and store at -80°C	Store isolate once grown in 2 Microbank vials and keep at -80°C
		Study Tube 2: Sample for storage	General CSF storage: Unspun sample	Store at -80°C
			India ink storage: centrifuge 500µl of CSF at 2000g for 10 min and store the pellet at -80°C.	
			qPCR sub-study: centrifuge all remaining CSF at 2000g for 10min and store supernatant and pellet in different cryovials at -80°C	
	EDTA (4ml) x 3	Monitoring Tube 1: FBC	After FBC, spin at 800 <sup>x</sup> g for 15min and store remaining plasma as below	If being performed in research laboratory.
		Study Tube 1: Spin tube at 800 <sup>x</sup> g for 15min and remove plasma	Divide plasma into 2x2ml cryovials. Remove buffy coat and put into 1 x 2ml cryovial	Store at -80°C
		Study Tube 2: qPCR sub-study	Remove 1.2ml and store in 2ml cryovial containing ceramic beads; centrifuge remaining 3.8ml at 2000g for 10min and store plasma in 1.5ml Eppendorf tube	
	Yellow/Red Top Tube (4ml)	Monitoring Tube 1: ALT, urea, creatinine, and electrolytes	Spin and store remaining serum.	If being performed in research laboratory.
Day 28	EDTA (4ml)	Monitoring Tube 1: FBC	After FBC, spin at 800 <sup>x</sup> g for 15min and store remaining plasma at -80°C	If being performed in research laboratory.
	Yellow/Red Top Tube (4ml)	Monitoring Tube 1: ALT, urea, creatinine, and electrolytes	Spin and store remaining serum at -80°C.	If being performed in research laboratory.



Table 4: Shipments

Specimen	Destination	Shipment Date	Special Notes
CSF			
CSF – PK/PD Study			
Crypto Isolates			
CSF and Blood qPCR			
PK Plasma			
Serum			
Whole Blood			
Plasma			
Buffy Coat			
CSF and Blood PAXgene tubes			
PBMCs			

#### Training

Each staff member receives or has direct access to applicable Working Practice Documents (WPDs).

AMBITION-cm

Therapy Induction

Each staff member reviews the applicable WPDs once a year.

All WPD training is documented and tracked in the training log located in the Investigator Site File (ISF)

New staff is trained on applicable WPDs within 30 days of employment and all WPDs within 90 days of employment.

Staff members whose duties fall within this WPD scope are retrained within 14 days of the approval of each WPD revision.

Staff signatures: (signing below indicates that you have read this SOP and understand the material contained in it)

Date	Name (Please print)	Signature