

DOC NO.: FDA/CTD/FOR - 05

Ver. No.: 03

Effective Date: 04/02/2020

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TITLE: FOOD AND DRUGS AUTHORITY CLINICAL TRIALS QUARTERLY PROGRESS REPORT FORM

SECTION A: ADMINISTRATIVE INFORMATION				
FOOD AND DRUGS	Expected Date of	Actual Date(s) of	Protocol Number:	
AUTHORITY Clinical	Commencement (as	Commencement (at the Trial		
Trial Certificate	indicated on the	Centre(s):		
Number:	certificate):			
		//		
Trial Title:				
Trial Site(s)				
Reporting Period	P			
Dringing 1 Investigator		to	•••••	
Principal Investigator:	Name:			
	Address:	P	Phone:	
		Ν	Iobile:	
		E	E-mail:	
Co-Investigators:	Name(s):		hone:	
		Ν	Iobile:	
			E-mail:	
Other Trial Contact (if	Name:		hone:	
applicable):	Address:		Iobile:	
		E	E-mail:	
SECTION B: TRIAL STATUS (Check one category only)				
Enrolment h	has not begun			
Actively enrolling participants				
Enrolment closed on: (insert date): participants are receiving treatment/intervention				
Enrolment closed on: (insert date): participants are in follow-up only.				
Analyzing data				
Data analysis completed				



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SECTION C: INFORMATION ON PARTICIPANTS & TRIAL ACTIVITIES

a.	Number of persons consented				
b.	Number of persons screened				
c.	c. Number of persons consented and screened who are eligible for the trial				
d.	d. Number of participants to which the investigational product(s) has been administered				
e.	e. Number of participants left to be enrolled into the trial				
f.	Number of participants who have discontinued the trial:by Investigator:				
	• voluntarily:				
	• due to SAE:				
	• lost-to-follow-up:				
g.	Have there been any Serious Adverse Events (SAEs)?	Yes No			
h.	Total number of SAEs: (attach line list of SAEs documented for the quarter)				
i.	Have these SAEs been reported to the Food and Drugs Authority	Yes No			
j.	If No, explain				
k.	Have there been any changes to the protocol since the Food and Drugs Authority approved?	Yes No			
1.	Is this amendment submitted to the Food and Drugs Authority?	Yes No			
m.	If No, explain				
n.	Date for the end of the trial				
0.	Date for the final trial report				



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SECTION D: COMMENTS (if any)

SECTION E: SIGNATURE

Signature of Principal Investigator

Date

Return this form and all supporting documentation to: THE CHIEF EXECUTIVE FOOD AND DRUGS AUTHORITY P. O. BOX CT 2783, CANTONMENTS, ACCRA or submit via e-mail to <u>clinicaltrials@fda.gov.gh</u>