			DISEASE	Investigational Products (IPs)/IP	DATE OF RECEIPT OF	PRINCIPAL			STATUS & DURATION OF	
N/O 1	KANGAROO CARE	Phase IV	Low Birth Weight	Peer Support Baby Box containing clothing kits and Bempu	APPLICATION  20th March 2025	INVESTIGATOR  Dr. Adziri Sackey	1. Korle-Bu Teaching Hospital (KBTH) 2. Sunyani Teaching Hospital	APPLICANT Center for Learning and Child Development (CLCD)	Application Approved, 21 months.	PURPOSE/AIM OF STUDY  1.Evaluate the effectiveness of peer support and the Baby Box in increasing KC coverage at home. 2. Examine contextual factors associated with intervention effectiveness using mixed methods.
				Polypropylene Mesh Medical device			1. Lawra District Hospital 2. Debiso District Hospital 3. St. Martins De Pores Hospital, Elikwe 4. Holy Family Hospital – Berekum 5. Holy Family Hospital – Techiman 6. Twifo-Praeso Government Hospita 7. Saltpond Government Hospital 8. Salaga Municipal Hospital 9. Sandema District Hospital 10. War Memorial Hospital 10. War Memorial Hospital 11. Bongo District Hospital 12. Begoro District Hospital 13. Sefwi-Wiswoo District Hospital 14. Baabiani District Hospital 15. St. Peter's Hospital, Nadoso 16. SDA Hospital, Agona Wiemose 17. SDA Hospital, Kwadsos 17. SDA Hospital, Kwadsos 17. SDA Hospital, Kwadsos	University of Birmigham, Dr. Birgit Whitman. Research Governance	Application Approved, 27	Primary objective: To assess if medical practitioners (MPs) can effectively perform mesh inguinal hemia repair compared to fully trained surgeons in adult patients with non-complicated inguinal hemia.  Secondary objectives:  To compare the impact of the intervention on:  O Surgical site infection and reoperation rates at 30 and 90 days after surgery or Recurrence at 90 days and one year after surgery.  Hemia-specific quality of life one year after surgery.  Change in quality of life from before to after surgery.  O Chorality of life from before to after surgery.  O Intervention of surgery.  Duration of surgery.  To explore the applicability of the trial's results by assessing the proportion of MPs requiring assistance from fully trained surgeons during inguinal hemia repairs.
	TIGER	Phase III	Surgery	Doxycycline, Moxidectin, and Albendazole Allopathic Drug	18th June 2024	Prof. Stephen Tabiri  Prof. Alexander Yaw	Kumasi Central Collaboration	Team  Kumasi Central Collaboration	months  Application Approved, 39	overall surgery costs.  Primary Objectives: To assess the effectiveness of the respective treatment regimens doxycycline (DOX), moxidectin +albendazole (MoxA) or standard mass drug administration (MDA) by comparing the proportions of the at baseline Slicine "Fliarial test Strip (FTS) positive participants who were included in the trial (eligible participants) and who became FTS-negative at 24 months after treatment onset. For all objectives: follow-ups for untreated participants will be based on the schedule of
	TAKE OFF T&T SEMAGLUTIDE	Bioavailability	Filariasis	Oral  Semaglutide sublingual tablets Allopathic Oral	21st August 2024  30th December 2024	Prof. George Obeng Adjei	Research  Azidus Laboratories Tema Freezone	Research  GFC Pharma LLC	Application Approved 4 months	the assigned treatment group of the community  To evaluate the bioavailability of Semaglutide sublingual tablets 1 mg following oral (Sublingual) administration in healthy subjects under fasting condition.
	SMAART MAP (new); Renal function domain			Paracetamol Allopathic drug Rectal/Oral/Nasogastric	28th March 2024/24th May 2024	Professor Daniel Ansong	Komfo Anokye Teaching Hospital Department of Child Health, Kwame Nkrumah University of Science and Technology		Application approved, 27 months	PRIMARY OBJECTIVE  Our primary objective is to test whether regularly dosed paracetamol given over 66 hours (corresponding to 72 hours exposure) will reduce levels of creatinine in children at high risk of renal impairment compared to standard of care; thus determining if paracetamol can reduce the evolution of kidney injury in severe malaria. SECONDARY OBJECTIVES  AND OBJECTIVES  OF THE OBJEC
6	SMAART MAP (new); Anaemia domain	Phase III	Anaemia	Whole Blood and Packed Blood Cells Transfusion	28th March 2024/24th May 2024	Professor Daniel Ansong	Komfo Anokye Teaching Hospital Department of Child Health, Kwame Nkrumah University of Science and Technology	Imperial College London	Application approved, 27 Months	PRIMARY OBJECTIVE  Our primary objective is to test whether giving a whole blood transfusion compared to red cell concentrates in children with severe malaria and severe anaemia leads to improved haemoglobin recovery and reduces the need for secondary transfusions.  SECONDARY OBJECTIVE  Our secondary objective is to assess the impact of whole blood vs red cell concentrate transfusions on other clinical outcomes such as mortality and readmission at 90 days and to understand the safety profile of both types of transfusions further by comparing grade 3 and 4 adverse events (AEs) and AEs of any grade related to the transfusions.

SMAART MAP (new) Cerebral malaria 7 domain	; Phase III	Cerebral malaria	Levetiracetam Allopathic drug Intravenous	28th March 2024/24th May 2024	Professor Daniel Ansong	Komfo Anokye Teaching Hospital Department of Child Health, Kwame Nkrumah University of Science and Technology	Imperial College London	Application approved, 27 Months	PRIMARY OBJECTIVE(S) Our primary objective is to test whether that leveliracetam given to children with seizures in their current episode of malaria but prior to admission will help prevent further seizures.  SECONDARY OBJECTIVE(S) Objective is to assess the impact of levetiracetam on other outcomes including mortality and readmission at 90 days and to investigate its safety profile in this patient population by grade 3 and 4 adverse events (AEs), solicited AEs, and AEs of any grade related to anticonvulsants.  •An additional objective is, where it is possible, to store blood spots on filter papers, in order to further assess the pharmacokinetics of levetfracetam in this patient population.
8 SHINE-1	Phase Ⅲ	Human Papilloma Virus (HPV)	Innovax 9 (Recombinant Human Papillomavirus 9-valent Vaccine (Escherichia Coli) Vaccine Intramuscular	3rd July 2024	Dr. Nana Akosua Ansah	Navrongo Health Research Center (NHRC)	РАТН	Application approved, 32 months	Primary Objective: *To evaluate NI of immune response for the Innovax 9vHPV vaccine administered in a single-dose schedule to that of Gardasil 9 against oncogenic HPV types (HPV-16, -18, -31, -33, -45, -52, and -58) in healthy girls 9– 14 years of age, 24 months after vaccination.  *To evaluate NI of immune response for the Innovax 9vHPV vaccine administered in a single-dose schedule to that of Gardasil 9 against oncogenic HPV types (HPV-16, -18, -31, -33, -45, -82, and -58) in healthy young women 15–20 years of age, 24 months following vaccination.  *Secondary Objective (Immunogenicity) To evaluate NI of immune response for the Innovax 9vHPV vaccine administered in a single-dose schedule to that of Gardasil 9 against HPV types 6 and 11, 24 months following vaccination
9 URIB-PAP	Phase I	Human Papilloma Virus (HPV)	Urine collection device for HPV testing Medical device Intravaginal	20th June 2024		Korle-Bu Teaching Hospital (KBTH)	University of Michigan Department of Obstetrics and Gymecology	Application approved, 11 months,	Aim(s)  *To explore the acceptability and feasibility of our device among KBTH healthcare clinicians.  *To validate that our device facilitates highly accurate urine-based HPV screening.  *To explore the acceptability and feasibility of our device among KBTH patients.  *Specific objectives  *Examine clinician acceptability of our device.  *Examine clinician perspectives on the feasibility of utilizing our device to screen patients.  *Compare detection rates of HPV for our device versus Pap smears.  *Examine patient satisfaction with our device versus Pap smears.  *Understand patient experiences, perspectives, and attitudes regarding HPV screening.
AZIDUS 10 BUPRENORPHINE	Bioequivalen ce Study	Analgesic	Buprenorphine Allopathic Drug Oral	30th July 2024	Dr. George Obeng Adjei	Azidus Laboratories Tema Freezone	Wes Pharma Inc,USA	Application approved, 2 months	Primary Objective(s): The objective of this pilot study is to evaluate the Test formulation in comparison to the Reference Standard and to generate pharmacokinetic data that can be used to design a pivotal bioequivalence study
PMC RTSS SUB 11 STUDY	Phase III	Malaria	Sulphadoxine/Pyrimethamine + Amodiaquine, Sulphadoxine/Pyrimethamine, RTS,S/AS01E Vaccine Allopathic drug and Vaccine Oral and intramuscular injection	8th May 2023	1.Dr. Dennis Adu-Gyasi 2. Fr. Kwaku Poku Asante	Kintampo Health Research Center	Kintampo Health Research Center	Application Approved, 40 months	Primary objective The primary objective of the study is determination of whether children who have received PMC with SP or SPAQ together with the RTS,SIAS01E vaccine have lower levels of naturally acquired immunity to malaria, as measured by antibodies to blood stage malaria antigens, than children who have received the malaria vaccine alone when they reach the ages of 18 and 24 months of age, the age at which they cease to be eligible to receive PMC.  Secondary objectives of the study include -  1. Determination of whether children who have received PMC with SP or SP+AQ together with the RTS,SIAS01E vaccine have lower titres of anti-CSP antibody than children who have received the malaria vaccine alone at 10 months of age (one month after they have received a booster dose of the vaccine), at 19 months of age, (one month after they have received a booster dose of vaccine), and when they reach the age of 24 months.  2. Determination of whether children who have received PMC with SP or SPAQ together with the RTS,SIAS01E malaria vaccine have lower cellular immune responses to the CSP protein than children who have received RTS,SIAS01E alone when they reach the ages of 18 and 24 months.  3. Determination of whether the immune response to priming and booster doses of the RTS,SIAS01E vaccine is influenced by the presence of asymptomatic malaria parasitaemia at the time of vaccination.

12 REALISE	Phase III	Soil-Transmitte d Helminth Infections	Albendazole-Ivermectin Allopathic drug Oral	9th May 2024	Dr. Abraham Rexford Oduro Dr. Joseph Kwadwo Opare	Nzema East District, Western Region	Laboratorios Liconsa SA	Application Approved, 3 years	Primary objective 1. To evaluate and compare the safety of the FDC against ALB via mass drug administration (MDA). Secondary objective 1. To evaluate the effectiveness of one round of MDA with FDC compared to ALB against Trichurs trichura. Exploratory objectives 1. To evaluate the effectiveness of one round of MDA with FDC compared to ALB against Strongyloides stercorals by serology. 2. To evaluate the effectiveness of one round of MDA with FDC compared to ALB against Strongyloides stercorals by serology. 3. To evaluate the effectiveness of one round of MDA with FDC compared to ALB against Ascaris lumbircoides. 4. Describe the frequency of scables before and after the intervention in the two treatment arms. 5. To implement genomic surveillance as a tool to evaluate MDA effectiveness and monitor drug resistance emergence in T. trichiura. 6. To assess the role of the gut microbiome on the effectiveness of one round of MDA with ALB and FDC.
13 IMBRAVE 152	Phase III	Liver Cancer	Atezolizumab/Bivacizumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Tiragolumab/Ti	15th November 2023	Dr. Edward Amankwah Frimpong     Z. Dr. Asare Offei	Korle-Bu Teaching Hospital (KBTH)     Sweeden Ghana Medical Centre	F. Hoffmann-La Roche Ltd	Application Approved, 2 years 8 months	Primary Objectives:  *To evaluate the efficacy of atezolizumab plus bevacizumab plus tiragolumab compared with atezolizumab plus bevacizumab  Secondary Objectives:  *To evaluate the efficacy of atezolizumab plus bevacizumab plus tiragolumab compared with atezolizumab plus bevacizumab  *To evaluate the safety of atezolizumab plus bevacizumab plus tiragolumab compared with atezolizumab plus bevacizumab plus tiragolumab compared with atezolizumab plus bevacizumab plus tiragolumab  *To characterize the PK profile of atezolizumab plus bevacizumab plus tiragolumab  *To evaluate the immune response to tiragolumab and atezolizumab
14 NANOX.ARC		Radiographic	Nanox.ARC Medical device NA	11th March 2024	Dr. George Boateng KYEI	University of Ghana Medical Centre	NANO-X MAGING LTD	Application Approved, 2 years	Primary Objective:  *To assess safety and clinical performance of Nanox.ARC DTS in providing additional information to conventional 2D radiography when evaluating adult individuals with known or suspected radiographic abnormalities.  Secondary  Objectives  *To evaluate the ability of Nanox.ARC DTS to reduce the need for a CT/MRI or other advanced imaging modality  *To evaluate the ability of Nanox.ARC DTS to increase the level of confidence of the reader in identifying/excluding an abnormality.  *To evaluate physician reading time of Nanox.ARC DTS compared to CT/MRI or other advanced imaging modality  *To evaluate the length and extent of the learning curve of reading the tomosynthesis images  Safety  Objectives  Safety  Objectives  Safety objective is to collect safety information, including type and number of adverse events, serious adverse events, and device issues.
15 REVIVE	Phase III	Advanced HIV	Zithrolide (Azithromycin) Allopathic drug Oral	14th March 2024	Dr. Yasmine Oladele I. Hardy Prof. Daniel Ansong	Kumasi (Bantama, Suntreso and Atonsu)	Hamilton Health Sciences through its Population		Primary Objective: The primary objective is to determine whether azilthromycin is an effective and safe intervention to reduce excess mortality in adults with advanced HIV (CD4 ≤ 100 cells/mm3).  Secondary Objective: Secondary objectives include exploring effects on mortality and hospitalisation at early and late timepotins, impact on incident infection, and cause of death.
MICRONUTRIENT 16 SUPPLEMENTATION	Phase III		Micronutrient (Effervescent powder; Orange flavored; Contains multiple vitamins and minerals) Food supplement Oral	15th April 2024	Prof. Francis Bruno Zotor	University of Health and Allied Sciences	InnoNext Sårl	Application Approved, 3 years 8 months	The primary objective of the study to determine if micronutrient supplement improves the vitamin D status of the study participants with or without additional Nutrition Training and Healthy Lifestyle Coaching (herein referred to as NuTHLIC). Vitamin D status will be assessed as serum 25(0H) D in serum. The secondary objectives of the study are to: 1. Determine if micronutrient supplementation improves the status of vitamin B12, zinc, magnesium and iron of the study participants that will receive a micronutrient supplement with or without Additional nutrition Training and Healthy Lifestyle Coaching (herein referred to as NuTHLIC). The nutrient status will be assessed as serum vitamin B12, serum rainic, serum magnesium, serum ferrith and RBC H. Batus through the assessment of the nutrient biomarkers as per point 1.  2. Assess the effectiveness of the micronutrient supplement with or without additional NuTHLIC on the flestyle habits and overall wellbeing through targeted questionnaires as assessed by theparticipants.

	MALHELMINTH 17 STUDY	Phase IV	Helminths infection/Malari a	Sulphadoxine-pyrimethamine and Amodiaquine - (SPAQ), Albendazole (ALB), Praziquantel (PZQ)/Allopathic drug Allopathic drug Oral	29th December 2023	Dr Muhammed Afolabi     Dr Kwaku Poku Asante		London School of Hygiene & Tropical Medicine	Application Approved, 13 months	Aim:  To evaluate the effectiveness and cost-effectiveness of integrating mass drug administration for helminth control with seasonal malaria chemoprevention in Ghanaian children Objectives:  Evaluate the effectiveness of combining SMC and deworming drugs in reducing the prevalence of anaemia and the intensity of malaria-helminth co-infections among a population of pre-school and school age children resident in a high burden country.  Determine the cost and cost-effectiveness of delivering an integrated malaria-dewormingapproach to the children.
	18 KALUMA STUDY	Phase III	Malaria	KLU156	27th October, 2023	Dr. Samuel Harrison     Dr. Patrick Odum Ansah	1. KHRC 2.NHRC	Novartis Pharma AG	Application Approved, 3years 9 months	Purpose This study aims to confirm the efficacy, safety and tolerability of KLU156, a fixed dose combination of ganaplacide (KAF156) and a solid dispersion formulation of lumefantine (SDF), when administered once daily for three days in adults and children ≥ 5 kg body weight and ≥ 2 months of age suffering from uncomplicated P, falciparum malaria (with or without other Pissmodium spp. co-infection). In the Extension phase, the safety, tolerability and efficacy of repeated treatment with KLU156 will be assessed for a maximum of two years in patients who did not experience any study treatment-related SAE (Serious Adverse Event) previously and who gave informed consent to participate in the Extension phase.
	SOY PEPTIDE 19 STUDY	Phase II	Malnutrition in cancer patient	Soy Protein Peptide Supplements (Vegalbum Supplement ) Food supplements Oral	10th February 2023	Prof. Christiana Nsiah- Asamoah	Cape Coast Teaching Hospital (CCTH)	South China University of Technology	Application Approved, 12 months	Objective: The aims of this study are evaluate the efficacy of food-borne (soybean) peptides in reducing mainutrition in cancer patients and (2) the secondary objective is to assess the impact of the peptides on hemoglobin levels, kidney function, liver function, and C-reactive protein levels in cancer patients.
	20 IAVI C105 STUDY	Phase II	Lassa Fever Disease	rVSVΔG-LASV-GPC Vaccine Vaccine Intramuscular Administration	7th August 2023	Prof. Kwadwo Koram	Noguchi Memorial Institute for Medical Research	International AIDS Vaccine Initiative (IAVI)/ Susan Adu- Amankwah	Application Approved/4 years 3months	Safety  *To evaluate the safety and tolerability of the rVSVΔG-LASV-GPC vaccine at 2 different dosage levels in adults, including PLWH, and in children. Immunogenicity  *To determine binding LASV-GPCspecific antibody responses induced by rVSVAG-LASV-GPC vaccine  *To determine neutralizing LASV-GPCspecific antibody responses induced by rVSVAG-LASV-GPC vaccine in a subset of participants in each group
:	VERTEX Trial-BANK 11 HOSPITAL	Phase IVIII	Kidney Disease	Inaxaplin (VX-147) Allopathic drug Oral	22nd November 2023	Dr. Charlotte Osafo	The Bank Hospital		Application Approved 4 years	Primary objectives evaluate the efficacy of VX-147 to reduce proteinuria the efficacy of VX-147 no renal function as measured by eGFR slope Secondary objectives vealuate the efficacy of VX-147 to decrease the risk of the composite clinical outcome 'To evaluate the safety and tolerability of VX-147 identify the optimal dose from Phase 2 to carry forward to Phase 3 characterize the plasma pharmacokinetics (PK) of VX-147
:	22 ROBOCOW	Phase II	Postoperative Respiratory Tract Infections in abdominal surgery	0.2% Chlorhexidine Digliconate Mouthwash Oral	10th January 2023	Dr. Mohammed Sheriff		Dr. Mohammed Sheriff	Application Approved 5 Months	Primary Objective determine whether perioperative use of 0.2% chlorhexidine mouth wash reduces the rate of postoperative respiratory tract infections in 30 days postoperative period compared to placebo among patients undergoing midline laparotomy.  1.To assess the impact of the intervention on 30-day postoperative mortality 2.To determine the impact of the intervention on length of hospital stay 3.To determine whether the intervention impacts on the 30-day unplanned readmission rates due to a respiratory complication 4.To assess the effect of the intervention on time to return to normal activities
	23 INTS GMMA STUDY	Phase II	Typhoid	GVGH INTS-GMMA vaccine (GSK4077164A) Vaccine Intramuscular injection	17th May 2023	Professor Ellis Owusu- Dabo	KNUST-IVI Collaborative Centre	GlaxoSmithKline Biologicals SA	Application Approved, 3 years 4 months	To identify the preferred dose of each component of the iNTS-GMMA vaccine (Dose A [low], Dose B [medium], or Dose C [high]) for infant participants 6 weeks of age     Z. To evaluate the safety and reactogenicity of the iNTS-GMMA vaccine in all participants

24	VERTEX Trial-KBTH	Phase II/III	Kidney Disease	Inaxaplin (VX-147) Allopathic drug Oral	8th May 2023	Dr. Dwomoa Adu	Korle-Bu Teaching Hospital (KBTH)	Vertex Pharmaceuticals Incorporated	Application Approved 4 years	Primary objectives evaluate the efficacy of VX-147 to reduce proteinuria the efficacy of VX-147 on renal function as measured by eGFR slope Secondary objectives evaluate the efficacy of VX-147 to decrease the risk of the composite clinical outcome 'To evaluate the safety and tolerability of VX-147 identify the optimal dose from Phase 2 to carry forward to Phase 3 characterize the plasma pharmacokinetics (PK) of VX-147
25	FITBIT/XIAOMI	Phase III	Monitoring of Vitals in pediatric appendectomy and trauma patients	Fitbit Inspire 2, Xiaomi Mi Smart band 6 Medical device	20th March 2023	Dr. William Appeadu- Mensah	Korle-Bu Teaching Hospital (Paediatric Surgery Unit, Accident Centre)	1. Dr. Fizan Abdullah Ann and Robert H. Lurie Children's Hospital 2. Dr. Hassan Ghomrawi Northwestern University	Application Approved, 2 Months	Aim(s)  To establish the feasibility of a Fitibit/Xiaomi band-based wireless monitoring system for post-operative inpatient monitoring and monitoring of patients following trauma in the accident center. pecific objectives  The specific objectives of this study are to:  1. Determine the feasibility of implementing a band-based wireless monitoring system for post-operative, in-hospital monitoring of pediatric appendectomy patients, and for emergency department monitoring of pediatric and adult trauma patients.  2. Compare the vital signs recorded manually to those collected by wearable devices
26	PMC TRIAL	Phase III		Sulphadoxine/Pyrimethamine + Amodiaquine, Sulphadoxine/Pyrimethamine, RTS,S/AS01E Vaccine Allopathic drug and Vaccine Oral and intramuscular injection	8th May 2023	Dr. Kwaku Poku Asante	Kintampo Health Research Centre	PATH	Application Approved, 3 years 8 months	The primary objective is to determine the efficacy of the combination of RTS,S/AS01E and PMC with sulphadoxine/pyrimethamine alone (PMC SP) or RTS,S/AS01E and PMC with SP and amodiaquine (PMC-SPAQ) against clinical malaria among children up to 24 months of age compared with RTS,S/AS01E vaccine administered alone
27	PLATINUM	Phase Ila	Malaria	INE 963, Cipargamin (KAE609), KLU156/ KAF156/LUM-SDF, Coartem/Riamet Allopathic drug	29th March 2023	Dr. Patrick Odum Ansah	Navorongo Health Research Center (NHRC)     Kintampo Health Research Center (KHRC)	Novartis Pharma AG	Application Approved 21 Months	Part A: To assess the parasite clearance time (PCT) of oral doses of an antimalarial agent administered as monotherapy in patients with uncomplicated P. falciparum malaria Part B: To assess the effect on adjusted 28-day cure rate of an anti-malarial agent administered orally as combination therapy versus the standard of care (SoC) in patients with uncomplicated P. falciparum malaria
	NOVIC TRIAL	Phase III	Postpartum Hemorrhage (PPH)	Jada System (Intrauterine Vacuum Induced Hemorrhage Control Device) Medical device Vaginal	5th April 2022	Dr. Samuel A. Oppong	Korle-Bu Teaching Hospital (KBTH)     Korlend Anokye Teaching Hospotal     (KATH)		Application approved, 48	Study Objectives  1. To evaluate the effectiveness of the Jada® System, compared to standard care, in treating PPH, as measured by maternal survival without surgical intervention.  2. To assess the safety of the Jada® System, compared to standard care, in treating PPH, as measured by rate of composite adverse events potentially related to the device, including gental tract injury, uterine perforation or rupture and endometritis.  3. To estimate the cost-effectiveness of the Jada® System, compared to standard care, in treating PPH, as measured by incremental cost per quality-adjusted life year.
29	VERTEX Trial	Phase II/III	Kidney Disease	Inaxaplin (VX-147) Allopathic drug Oral	23rd December 2022	Professor Sampson Antwi	Komfo Anokye Teaching Hospital (KATH)		Application approved, 4 years	Primary objectives evaluate the efficacy of VX-147 to reduce proteinuria the efficacy of VX-147 or renal function as measured by eGFR slope Secondary objectives -To evaluate the efficacy of VX-147 to decrease the risk of the composite clinical outcome -To evaluate the safety and tolerability of VX-147 tientify the optimal dose from Phase 2 to carry floward to Phase 3 -To characterize the plasma pharmacokinetics (PK) of VX-147
30	COPE TRIAL	Phase III	Fistula	Healeanlo silicone lady Drain Valve menstrual Cup Medical device Intravaginal	2nd September 2022	Dr. Gabriel Y.K. Ganyaglo	Mercy Women's Catholic Hospital in Mankessim     Z Tamale Fistula Center in Tamale	Korle Bu Teaching Hospital	Application Approved, 15 Months	The aims of the study are to examine the effectiveness, comparative effectiveness, and acceptability of two vaginal menstrual cup models (cup and cup+) as a temporizing alternative to managing urinary leakage from vesico-vaginal fistula in both a clinical setting and a community setting, and to quantify non-surgical fistula management costs.

5	il PRAISE	Phase IVIII	Sickle Cell Disease	Oral FT-4-202 Pyruvate Kinase Activator and Placebo Allopathic drug Oral	2nd June 2022	1. Dr. Prince Agyapong - KHRC 2.Dr. Edeghonghon Olayemi - KBTH	Kintampo Health Research Center     Ghana Institute of Clinical Genetics,     KBTH	NOVO NORDISK COMPANY	Application Approved, 43 Months	Objectives of the study are: assess the efficacy of FT-4202 in adolescents and adults with SCD as compared to placebo as measured by improvement in hemoglobin (Hb) 2. To assess the efficacy of FT-4202 as compared to placebo on the annualized vaso-occlusive crisis (VOC) rate on clinical measures and sequelae of hemolysis 4. To evaluate the effects of FT-4202 on the sequelae of VOC assess changes in fatigue of sickle cell patients taking FT-4202
3	PROBIOTIC PILOT 12 STUDY	Pilot study	Malnutrition	Synbiotic (Nutraflora and Maltrin M100 P-95 and L. plantarum (Lp) and Placebo Food supplement	27th July, 2021	Dr Seyram Kaali	Kintampo Municipal Hospital	Dr. Kwaku Poku Asante	Application Approved 27 months	Primary A pilot trial to evaluate the administration of probiotic supplementation among pregnant women in the third trimester and effective colonization of the gut microbiome of their infants one-month post-partum. Secondary 1. To assess compliance of administering a synbiotic product (L. plantarum with Fructooligosaccharide) among pregnant women. 2. To assess birth outcomes among participants who receive synbiotic products compared to those on placebo. 3. To assess if maternal stool microbiome profoundly changes from immediately after childbirth to one-month post-partum. 4. To characterize the diversity of vaginal microbiomes among pregnant women in the study area. 5. To determine the safety of the probiotic supplementation among pregnant women from 5 to 6 months until up to two weeks post partum.
3	13 ASAAP	Phase III	Malaria	Arthemeter + Lumefantrine, Atovaquone //Proguanil Hydrochloride and Placebo (P- Dragees Rosa Lichtenstein)  Allopathic drug  Oral	4th October 2021	1. John Humphrey, AMUASI Dr Oumou Maiga Ascofare	St. Francis Xavier Hospital	Kumasi Centre for Collaborative Research (KCCR), Kumasi, Ghana	Application Approvedl 21 months	The overall aim of this phase III clinical trial(main study = study II) is to develop a readily deployable highly efficacious, safe and well tolerated antimalarial triple combination therapy for young children.  This is achieved by evaluating the efficacy, safety and tolerability of artemether-iumefantrine (AL) + atovaquone-proguanil (AP) tri-therapy (AL+AP) compared to standard AL therapy (+placebo) for the treatment of uncomplicated Plasmodium falciparum malaria in African children aged femonths to 10years.
3	14 PIVOT STUDY	Phase II	Sickle Cell Disease	Hydroxyurea and Placebo Allopathic drug Oral	18th June 2021	Dr. Yvonne A. Dei- Adomakoh     Dr.Catherine Segbefia	Korle-Bu Teaching Hospital	Cincinnati Children's Hospital Medical Center	Application Approved 5 years	To measure the toxicities of hydroxyurea treatment on laboratory parameters.  To assess the effects of hydroxyurea treatment on a variety of sickle-related clinical and laboratory parameters in a large cohort of children and adults with HbSC disease.  To identify which study endpoints are suitable for a future Phase III trial of patients with HbSC disease receiving hydroxyurea therapy.
3	5 RECOVERY	Phase III	Covid-19	Infliximab, Dexamethasone Allopathic drug Oral and/or Intravenous	21st May, 2021	Dr. John H. Amuasi	Komfo Anokye Teaching Hospital Ghana Infectious Disease Centre	University of Oxford Clinical Trials and ResearchGover nance.	Application Approved 2 years	For each pairwise comparison with the 'no additional treatment' arm, the primary objective is to provide reliable estimates of the effect of study treatments on all-cause mortality at 28 days after randomisation (with subsidiary analyses of cause of death and of death at various timepoints following discharge). The secondary objectives are to assess the effects of study treatments on duration of hospital stay; and, among patients not on invasive mechanical ventilation at baseline, the composite endpoint of death or need for invasive mechanical ventilation or ECMO.
3	TyVEGHA 6	Phase IV	Typhoid fever	Vi polysaccharide-tetanus toxoid conjugate vaccine (Vi-TT), Meningococcal Group A conjugate vaccine (MCV-A 5) Vaccine Intramuscular	9th April 2021	Prof. Ellis Owusu-Dabo	Agogo Trial Center/KNUST- International Vaccine Institute (IVI) Collaborating Center	International Vaccine Institute	Application Approved Study commenced 3 Years 5 months	The purpose of the study is to determine the total protection conferred by single-dose vaccination with Vi-TT against blood culture-confirmed symptomatic S. Typhi infection in the intervention vaccine clusters, compared with the control vaccine clusters  • To investigate the safety outcomes associated with Vi-TT vaccination in the intervention vaccine recipients compared with the comparator vaccine recipients. To determine the overal protection of Vi-TT vaccination against blood culture-confirmed symptomatic infection caused by S. Typhi in intervention clusters compared with control clusters  • To determine the total protection of Vi-TT vaccination against severe TF in the intervention vaccine recipients compared with the comparator vaccine recipients  • To determine the overall protection of Vi-TT vaccination against severe TF caused by S. Typhi in intervention clusters compared with control clusters  • To investigate the total protection of Vi-TT vaccination against clinical TF (defined below in Trial Outcome Measures') in the intervention vaccine recipients compared with the comparator vaccine recipients in Trial Cultored with control clusters  • To investigate the overall protection of Vi-TT vaccination against clinical TF in intervention clusters compared with control clusters  • To measure the indirect protection conferred by single-dose vaccination with Vi-TT against blood culture-confirmed symptomatic S. Typhi infection in the intervention vaccine clusters.

37 FILOVIRUS	JS STUDY F	Phase I/II/III	Filovirus disease	Vaccine and therapeutic candidates Vaccine and therapeutic Intramuscularand oral	1st April 2025	Dr John Amuasi/Prof George Kyei-Boateng	KCCR Noguchi Memorial Institute for Medical Research		Application Pending Approval	Primary objectives To determine the reactogenicity and safety of candidate filovirus vaccine(s) among healthy volunteers. To determine the immunogenicity of the candidate filovirus vaccine(s). Secondary Objectives To determine the durability of filovirus-specific induced immune responses following vaccination. To identify factors influencing vaccine-induced immune responses among that participants.
38 CLARITY A	AFRICA F	Phase III	Stroke	Cilostazole/Aliopathic/Oral	11th June 2025	Prof. Fred Stephen Sarfo		Prof. Akwasi Antwi Kusi/Prof. Fred Stephen Sarfo	Application Pending Approval, 5 years	Study Goal:  Overall objective of the CiLostAzol for pRevention of recurrent sTroke in Africa (CLARITY-AFRICA) study is to deploy a hybrid study design to demonstrate the efficacy & safety of cilostazol twice daily in reducing MACE over 24 months vs. placebo among 1,100 recent stoke patients encountered at 12 hospitals in Ghana.  Secondly, CLARITY-AFRICA also seeks to develop an implementation strategy for routine integration & policy adoption of cilostazol for post-stroke cardiovascular risk reduction in an under-resourced system.
39 SPARKLE	E F	Phase III	Sickle Cell Disease	Crizanlizumab/Allopathic/Oral	9th May 2025	Dr Lucy Osei Ababio     Prof Yvonne Dei- Adomakoh     Dr Frank Baiden     Dr Tsiri Agbenyega     Dr Lesey Osei	1.Navrongo Health Research Centre     2. Ghana Institute of Clinical Genetics     Korfe Bu Teaching Hospital 3.     University of Health and Allied     Sciences     4. Agogo Presbyterian Hospital     5. Directorate of Child Health Komfo     Anokye Teaching Hospital	Novartis Pharma AG	Application Pending Approval, 5 years	Primary Objectives  To compare the efficacy of 5 mg/kg of crizanlizumab versus placebo, with or without hydroxyurea/hydroxycarbamide, on the annualized rate of VOCs* that are HCP managed (including VOCs leading to management at a health care facility or those managed via remote consultation) over the planned 52-week treatment period in SCD patients aged 12 years and older with a history of frequent VOCs (4-12 events in 12 months prior to the screening visit)
40 FLORAL S	STUDY F	Phase III	Sickle cell disease	Etavopivat Allopathic Oral	27th January 2025	Dr Seyram Kaali     Dr Edeghonghon Olayemi	Chana Institute of Clinical Genetics	Novo Nordisk A/S	Application Pending Approval, 61 monhs	Primary objectives:  To investigate long-term safety of etavopivat in adults, adolescents and children with SCD, SCDTD, TDT or NTDT transferring from other studies with etavopivat Secondary objectives: To investigate long-term clinical efficacy measures of etavopivat treatment in adults, adolescents and children with SCD transferring from other studies with etavopivat. To evaluate the effects of etavopivat on hospitalisations in adults, adolescents and children with SCD transferring from other studies with etavopivat.
41 NEOSEP 1	1 F	Phase III/IV	Neonatal Sepsis	1.Fomicyt 2. Flumarin Allopathic Oral	14th January 2025	Dr. John Humphrey Amuasi		Global Antibiotic Research & Development Partnership (GARDP)	Application Pending Approval,	Part 1 objectives & interventions: The purpose of Part 1 is to confirm that the recommended doses of fosfomycin and flomoxef, when used in combination with each other or with amrikacin to be studied in Part 2, will provide adequate drug exposure in neonates with sepsis. A secondary objective is to collect safety data.  Part 2 objectives & interventions: The purpose of Part 2 is to provide a ranking of eight different clinically relevant antibiotic regimens for first-line empiric and second-line (after lack of response/deterioration) treatment in terms of 28-day mortality as the primary outcome measure. It will flexibly compare these multiple different relevant treatment regimens to enable the trial to be run in sites worldwide with very different background rates of resistance anable the trial to be furning clinical care by randomising each participant to locally relevant antibiotic regimens agreed prior to site initiation.
42 HIBISCUS		Phase III	Sickle cell	Etavopivat Allopatric Oral	26th November 2024	Dr Seyram Kaali     Dr Patrick Ansah	Kintampo Health Research Center	Novo Nordisk		Primary Objective  demonstrate superiority of treatment with etavopivat versus placebo in adolescents and adults with SCD Objectives  Objectives  To evaluate clinical efficacy measures of etavopivat treatment versus placebo in adolescents and adults with SCD  To evaluate clinical efficacy measures of etavopivat treatment versus placebo in adolescents and adults with SCD  "To assess clinical efficacy measures of etavopivat treatment versus placebo in adolescents and adults with SCD  To assess clinically meaningful improvement in fatigue and functional exercise capacity and QOL measures of adolescents and adults with SCD taking etavopivat treatment compared to placebob

						Tp			
13 BILI-RULER		Neonatal Jaundice	Bilicare     Bilicare     Bilicare Medical Device	25th November 2024	Dr Kwaku Poku Asante	Kintampo Health Research Centre	Bill & Melinda Gates Foundation	Application Pending Approval,	Aims and Objectives  The objective of this substudy is to assess the ability of Bill-ruler used in community settings in identification of severe hyperbilirubinemia in neonates, as compared to visual assessment and TCB, among those born in the Pregnancy Risk, Infant Surveillance, and Measurement Alliance (PRISMA) Maternal and Newborn Health (MNH) Study. To achieve this aim, four statistical objectives were identified: 1. To estimate the level of agreement between Bill-ruler, visual assessment, and TCB values. 2. To estimate the level of agreement between Bill-ruler, visual assessment, and TCB among binary diagnostic categories (refer to a facility for treatment of hyperbillrubinemia' versus' do not refer to a facility of the state of th
14 ZERO POINT FIVE	Phase III	Hookworm infection, Ascaris lumbricoides, and Trichuris trichiura	ZP5-9676 Allopathic Drugs Oral	8th August 2024	Dr. Kwaku Poku Asante	Kintampo Health Research Centre (KHRC)	Zero Point Five Therapeutics	Application Pending Approval,	Primary objective:  • To evaluate the efficacy of ZP5-9676 for the treatment of hookworm (A. duodenale and N. americanus), Ascaris lumbricoides, and Trichuris trichlura in Participants between the ages of 6 months and 59 years.  Secondary objective:  • To evaluate the safety and tolerability of ZP5-9676 for the treatment of hookworm (A. duodenale and N. americanus), Ascaris lumbricoides, and Trichuris trichlura in Participants between the ages of 6 months and 59 years.
AZIDUS IS CEFUROXIME	Bioequivalen ce Study	Antibiotic	Cefuroxime Axetil Tablets Allopathic Drugs Oral	30th July 2024	Dr. George Obeng Adjei	Azidus Laboratories Tema Freezone	OA&J Pharmaceuticals Ltd	Application Pending Approval,	Primary Objective: To assess the bioequivalence between Test (T) and Comparator (R) formulations
AZIDUS 16 ACECLOFENAC	Bioequivalen ce Study	Analgesics	Aceclofenac tablets Allopathic Drugs Oral	30th July 2024	Dr. George Obeng Adjei	Azidus Laboratories Tema Freezone KBTH	OA&J Pharmaceuticals Ltd	Application Pending Approval,	Primary Objective(s):  • To evaluate and compare the relative bioavailability of two different test formulations (T1 & T2)  • To generate pharmacokinetic data that can be used to design a pivotal bioequivalence study
						KATH TTH Greater Accra Regional Hospital Sunyani Regional Hospital			
						Cape Coast Teaching Hospital Effla Nkwanta Regional Hospital Eastern Regional Hospital, Koforidua			Primary Objective:
			Drapes and Gown (Laparotomy drape)			Holy Family Hospital, Berekum  Holy Family Hospital, Techiman  Salaga Municipal Hospital  Goaso Municipal Hospital.			To assess whether reusable drapes and gowns are non-inferior in reducing SSI within 30 days of surgery compared to disposable (single-use) drapes and gowns.  Secondary Objective  - Assess the cost of using reusable versus disposable drapes and gowns  - Analyze the carbon footprint of reusable compared to disposable drapes and gowns:  - Investigate the rate of surgical site infections (SISs) associated with reusable versus
17 DRAGON	Phase I	Surgery	Medical Device	22nd July 2024	Prof. Stephen Tabiri	Ho Teaching Hospital St. Theresa Hospital	University of Birmingham	Application Pending Approval,	disposable drapes and gowns.  - Evaluate the patient experience of surgical site infections (SPECIES)

										AIM:  • To assess the effect of indispensable amino acids supplementation on
										environmental enteric dysfunction among children (18-36 months) with stunting.
										Specific objectives:
										Measure the effects of the indispensable amino acid supplementation on the change in child weight from baseline to end line.
						Dr.Regina Turkson				Determine the change in gut permeability due to IAA supplementation as assessed by L/R ratio.
						Dr.Charles Apprey				Determine the change in gut digestive capacity due to IAA supplementation as
				Amino Acid Mix (AA Mix)		Dr. Seyram Elom Achoribo		International		assessed by the 13C-sucrose breath test.  • Determine the change in plasma protein absorption index by Dual Stable isotope
AM	MINO ACID		Enteric Dysfunction/Nut	Food Supplement		Dr.Mame Yaa Adobea	Princess Marie Louise Children's	Atomic Energy Commission,		Technique (DSIT).  • Determine the changes in bacterial translocation, inflammation, damage, and peptide
48 SU	JPPLEMENTATION	Phase II	rition	Oral	10th July 2024	Nyarko,	Hospital (Accra)	Austria	Application Pending Approval,	transport in the gut
										To assess the performance of STANDARDTM Q hs- Malaria P.f/P.v Ag Test and
				Standard Q hs-Malaria Ag p.f/p.v&			NMIMR     Obom health center			STANDARDTM Q hs- Malaria P.f Ag Testin intended use settings for detecting P. falciparum and P. vivax infections in capillary and venous whole blood samples
				Standard Q hs-Malaria Ag p.f			3. Kofi Kwei CHPS compound,	SD		collected prospectively from patients with symptoms suggestive of malaria in
49 SD	D Biosensor MRDT	Phase III	Malaria	Medical device	2nd July 2024	Prof Linda Eva Amoah	Moree polyclinic,     Ewim Polyclinic	BIOSENSOR,IN C	Application Pending Approval,	accordance with the Technical Specifications Series for submission to WHO Prequalification – Diagnostic Assessment: Malaria rapid diagnostic tests.
										Buruli ulcer is a neglected disease caused by infection with Mycobacterium ulcerans (Mu), which manifests as large, disfiguring skin ulcers mainly in children aged 5 to 15
										years. Access to treatment in rural areas can be challenging and late presentation is
										typical, due to fear, stigma, suspicion about conventional medicine and economic consequences for poor families. The current recommended regimen of oral rifampicin
										together with intramuscular streptomycin or clarithromycin for 8 weeks is far from ideal, particularly given the increasing global threat of antimicrobial resistance. Although the
				Nitric oxide releasing gel, Vaseline Gauze						disease can be cured in most patients who adhere to this regimen, healing rates are highly variable even in patients with seemingly similar lesions.
				dressing materials			Kumasi Centre for Collaborative	Kumasi Center		
				Allopathic drug + medical device			Research in Tropical Medicine 2.Agogo Presbyterian Hospital	For Collaborative		The purpose of the study is to compare the healing measured by the percentage area reduction of EDX110 dressing with oral rifampicin and clarithromycin (EDX-RC) versus
BU 50	JRULINOX		Buruli Ulcer	Topical	24th September 2018	Prof. Richard Odame Phillips	Tepa Government Hospital     Dunkwa Government Hospital	Research (KCCR)	Study ended, Final Report yet to be submitted, 36 MONTHS	'Usual Care' with routine Vaseline gauze dressing and oral rifampicin and
00			Buruii Oicei	Торгові	24th September 2010	i inilips	т. данкwa Government поѕрна	(NOON)	be submitted, 30 MONTHS	clarithromycin (VG-RC).
				Rifampicin Capsules, Bacteria binding						
				dressing: acetate fabric coated dialkyl carbamoyl chloride (DACC)						
				Allopathic drug			•KCCR •Ga Fast munical hospital	London school of Hygiene and	Study and od Final Page 1 vet to	Compare the time to clearance of viable Mycobacterium from wounds of patients
							•Pakro Health Centre	Tropical	be submitted, 2 Years 6	treated with high-dose rifampicin and DACC dressings (HR-DACC) to those receiving
51 BU	JRULIRIFDAC	Phase III	Buruli Ulcer	Oral and Topical	12th December 2020	Prof. Richard Phillips	•Wassa Amenfi East Hospital	Medicine	Months	standard dose rifampicin and DACC dressings
										Main Aim
										This study explores the feasibility, acceptability, and outcomes of implementing sterile water injections (SWI) for the management of lower back pain among birthing women
										in Ghana.
										Specific Objectives
										Develop and deliver a training package for midwives on sterile water injections for managing lower back pain.
										Undertake implementation study in a tertiary hospital in Ghana to assess the
										feasibility and acceptability of implementing SWI for lower back pain.  3. Determine birth and neonatal outcomes of women with back pain who receive SWI
SW	VIS (STERILE	Feasibility	Lower Back	Sterile Water Injection				Dr. Jonas	Study ended, Final Report vet to	Explore the experiences of women who have had SWI for back pain in labour     Explore the experiences and perception of midwives and stakeholders regarding the
	ATER INJECTION)	study	Pain	Intradermal	6th December 2022	Prof. Sue Kruske	Korle-Bu Teaching Hospital (KBTH)	Awuku Afari	be submitted, 40 Months	implementation of SWI for managing back pain in labouring women.

53 A	CTIV TRIAL	Phase III	Covid-19	S-217622 Tablet and Placebo Allopathic drug Oral		1.Dr. Patrick Ansah 2. Dr. Seyram Kaali 3. Prof. Richard Odame Philips	Kumasi Centre for Collaborative Research (KCCR) 2. Kintampo Health Research Centr (KHRC) 3. Navrongo Health Research Centre	SHIONOGI INC.& Co Ltd		Primary Objective To determine if S-217622 will reduce the time to sustained symptom resolution through Day 29. Time to sustained symptom resolution is defined as the time from start of study intervention to the first day of 4 consecutive days with complete resolution of 13 COVID-19 symptoms on participant self-assessment AND alive and without hospitalization for any reason by Day 29. Hospitalization is defined as 224 hours of acute care, in a hospital or similar acute care facility, including emergency rooms, urgent care clinics, or facilities instituted to address medical needs of those with COVID-19. Secondary Objectives Key secondary objective: To determine the effect of S-217622 compared with placebo on the change from baseline in quantitative log10 SARS-CoV-2 RNA levels by PCR on NP swab at Day 4. Key secondary objective: To determine whether S-217622 reduces COVID-19 related hospitalization (adjudicated) and all deaths regardless of occurrence outside of hospital or during hospitalization (not adjudicated) through Day 29.
54 H	OPE KIDS 2	Phase III	Sickle Cell Disease	Voxelotor (GBT440) and Placebo Allopathic drug Oral	16th December 2020	Dr. Catherine Segbefia	*Korlebu Teaching Hospital Department of Child Health     *Sickle cell office Directorate Child(KATH)	Global Blood Therapeutics, inc	Study ended, Final Report yet to be submitted, 38 Months	The purpose is to evaluate the effect of voxelotor compared to placebo on the transcranial Doppler(TCD) time-averaged mean of the maximum velocity(TAMMV) arterial cerebral blood flow at 24 weeks in SCD participants >2 to < 15 years of age with conditional (170 to <200cm/sec) TCD flow velocity.
55 V	AT00008	Phase III	Covid-19	SARS-CoV2 prefusion Spike delta TM with AS03 adjuvant, monovalent, SARS-CoV2 prefusion Spike delta TM with AS03 adjuvant, bivalent, Matching placebo Vaccine Intramuscular			*Navrongo Health Research Centre *Kintampo Health Research Centre *Kwame Nkrumah University of Science and Technology (KNUST)	SANOFI	Study ended Final report yet to be submitted 41months 15days	To assess, in participants who are SARS-CoV-2 naïve, the clinical efficacy of the CoV2 preS dTM-AS03 vaccines for the prevention of symptomatic COVID-19 occurring ≥ 14 days after the second injection. To assess the safety of the CoV2 preS dTM-AS03 vaccines compared to placebo throughout the study.
	STAWOL	Phase II	Onchocerciasis /Filariasis	Rifampicin, Albendazole Allopathic drug Oral			-Bawku west -Builsa South -Nabdam Fumbisi -Garu-Tempane -Kayoro	Kumasi Centre for Collaborative Research (KCCR),	Study ended Final report yet to be submitted 24 months	The purpose of this study is to  *To show efficacy (Depletion of Wolbachia) of the combination of Rifampicin plus Abendazole against lymphatic filariasis using PCR compared to treatment with albendazole and "no treatment" (other than ivermectin) - Lymphatic Filariasis (LF) trial  *To show efficacy (depletion of Wolbachia and interruption of embryogenesis in female adult worms) of the combination of Rifampicin plus Albendazole, using PCR and immunohistology compared to treatment with albendazole and "no treatment" (other than ivermectin) - Onchocerciasis trial
C 57	ECOLIN	Phase III	Human Papiloma Virus (HPV)	Cecolin Vaccine Intramuscular	1st September 2020	Prof. Tsiri Agbenyega	•Agogo Asante Akim North District	PATH	Study ended Final report yet to be submitted, 30 months	The purpose of this study is to demonstrate the non-inferiority of Cecolin® administered on 0, 6-month; 0, 12-month; and 0, 24-month two-dose regimens, to Gardasi® using a 0, 6-month two-dose regimen, based on HPV Immunoglobulin G (IgG) antibody levels measured one month after the last dose for HPV types 16 and 18.
58 IU	IMO STUDY	Phase IV	Postpartum Hemorhage	Intrauterine Misoprostol and Sublingual Misoprostol/ Allopathic medicine			Department of Obstetrics and Gynaecology, Korle-Bu Teaching Hospital, Accra-Ghana.	Dr. Chidinma Peace Ohachenu	Study ended Final report yet to be submitted, 4 months	To evaluate the effectiveness of intrauterine misoprostol compared to sublingual misoprostol in the prevention of postpartum haemorrhage among women undergoing elective caesarean section in Korle-Bu Teaching hospital
A* 59	VAREF TV ROTA	Phase III	Gastroenteritis	1.Trivalent Rotavirus P2-VP8 Subunit Vaccine 2.Rotarix®/ Vaccine	9th April, 2019	1.Prof. George E. Armah 2.Dr. Alberta Amu	Dodowa Health Research Centre	РАТН		Diarrhea is the second-leading cause of death worldwide among children under the age of five, killing an estimated three quarters of a million children annually and hospitalizing millions more in developing countries. The most common cause of infantile diarrhoea is rotavirus and almost all children are infected by their third birthday regardless of geographical area or economic status. Infection is primarily via fecal oral route and improved sanitation alone will not control infection. Oral rotavirus vaccines have traditionally shown lower efficacy in Low and Middle Income Countries (LMICs) as compared to developed countries. Several theories proposed for this observation includes interference by other intestinal viruses or bacteria, neutralization of vaccine by maternally virus by maternally derived antibodies in breastmilk, etc. Some of these challenges may be obviated by a parenteral administered rotavirus vaccine. This study is therefore to demonstrate the efficacy and safety of the parenteral trivalent rotavirus vaccine in healthy infants (6a and <8 weeks old) to prevent severe rotavirus gastroenteritis compared with the orally approved Rotarix®

60	EBSI-LSV	Phase I	Lassa Fever	1.EBSI-LSV 2. Placebo/ Vaccine	1st September 2021	1.Dr Seyram Kaali 2.Dr.Patrick Ansah	Kintampo Health Research Centre     Navrongo Health Research Centre	Emergent BioSolutions (EBS)	Study ended Final report yet to be submitted 2 years	To evaluate the safety and tolerability of increasing dose levels of EBS-LASV vaccine administered as a single dose or two-dose series.     To evaluate the humoral immune response to EBS-LASV vaccine at various dose levels and dosing schedules for the purpose of selecting two regimens (dose and schedule) for further evaluation in a Phase 2 study.
61	SHEALIDO	Phase III	Rectal Examination	Optilube Active Sterile Lubricating Jetly     Shealube/ Lubricating gel	10th September 2020	Dr. Kekeli Kodjo Adanu	Ho Teaching Hospital	University of Health and Allied Sciences	Study ended Final report in the ICHE3 format yet to be submitted 12 months	This study is a randomized controlled trial which compares the effectiveness, complications and ease of use of shea butter as a surgical lubricant to lidocaine gel. The purpose is to:  10 determine the ease of use of shea butter by clinicians as compared to lidocaine gel as a lubricant for rectal examination.  10 determine the complication rate related to the use of shea butter as a lubricant for rectal examination.  10 ascertain the complication rate associated with the use of lidocaine gel as a lubricant for rectal examination.  10 compare the complication rate related to the use of shea butter to that of lidocaine gel.
62	INOVIO	1b	Lassa Fever	1.INO-4500 2.CELLECTRA™ 2000 3.SSC-0001/ Vaccine	30th September 2019	Prof. Kwadwo Ansah Koram	Noguchi Memorial Institute for Medical Research University of Ghana, Legon		Study ended Final report submitted 20 Months	The LASV DNA vaccine expressing the glycoprotein precursor (LASV GPC, Josiah strain matched) paired with intradermal EP is a promising vaccine platform that has been shown to elicit protective immunity and completely protect guinea pigs and non-human primates (NHP) against viremia, illness (acute and chronic), and death after Lassa virus exposure [26, 27] and protect NHPs from hearing loss (unpublished data). This LASV DNA vaccine, INO-4500, targets GPC because it represents the most conserved region in this genetically diverse virus. In the case of Lassa virus infection, the generation of a robust T cell response appears to be the key to protection from infection.  As such, the DNA-EP platform is highly amenable to this disease target. The purpose of this study is to evaluate the tolerability and safety of INO-4500 administered by ID injection followed by EP in healthy adult volunteers
63	MDGH-MOX	Phase I		Moxidectin tablet (2mg)/ Allopathic drug	February 2020	Dr. Nicholas Opoku	School of Public Health Research Centre, University of Health and Allied Health Sciences. Ho.	Medicines	Study ended Final report submitted. 12 months	To characterize the pharmacokinetics and safety of moxidectin in children (aged 4 to 11 years) and adolescents (aged 12 to 17 years) and to enable determination of an optimal dose for treatment of children 4 to 11 years
64	SPUTNIK LIGHT	Phase III	Covid-19	1.Sputnik Light Vector Vaccine 2.Placebo/	5th March 2021		Navrogo Health Research     Centre Dodowa Health Research     Centre Ghana		Study ended Final report yet to be submitted 8 months	The purpose of the study is to  - Assess efficacy of the Sputnik-Light vector vaccine against the SARS-CoV-2-induced coronavirus infection compared to placebo  - Assess tolerability and safety of the Sputnik-Light vector vaccine against the SARS-CoV-2-induced coronavirus infection compared to placebo  - Assess humoral immunogenicity of the Sputnik-Light vector vaccine against the SARS-CoV-2-induced coronavirus infection compared to placebo on Subset A.  - Assess protective properties of the Sputnik-Light vector vaccine against the SARS-CoV-2-induced coronavirus infection compared to placebo for prevention of serologically confirmed SARS-CoV-2 infection  - Assess efficacy of the Sputnik-Light vector vaccine against the SARS-CoV-2-induced coronavirus infection compared to placebo based on severity of COVID-19 disease
65	EMODEPSIDE	Phase II	Onchocerciasis	Emodepside (5mg) Allopathic drug	5th November, 2020	Dr. Nicholas Opoku	-School of Public Health Research Centre, (UHAS). -Municipal Hospital, Hohoe, Volta Region, Ghana -Kpassa, Nixwanta- North District, Oti Region, Ghana	DNDi (Drugs for Neglected Diseases initiative)	Study ended Final report yet to be submitted 67 months	The purpose of this study is to  *Ensure the safety and tolerability of emodepside after single oral doses administered as solution (liquid service formulation, LSF) or immediate release (IR) tablets in healthy male subjects  *Plasma PK of emodepside (solution and tablets), the effect of food on the bioavailability of emodepside

66	MAL 094	Phase IIb	Malaria	1.RTS,S/AS01E 2.Rables vaccine (Rabipur™)/ Vaccine	21st November 2016	Prof. Tsiri Agbenyega	Malaria Research Center, Agogo	GlaxoSmithKline Biologicals SA	be submitted	As part of GSK and PATH's commitment to develop a malaria vaccine for reduction of malaria disease burden in children and contribution to the malaria elimination goal, characterization of an optimal dosing regimen and boosting schedules are critical. Results of previous efficacy of stockedules are critical. Results of previous efficacy of stockedules with dose, and the preliminary results of MAL 071 study (recent controlled human malaria infection) were reviewed by the European Medicines Agency (EMA). There was evidence that demonstrated superior protection against malaria infection associated with the use of a fractional third dose in a 0, 1, 7-month schedule with a higher vaccine efficacy against malaria infection.  This study intends to establish Proof of Concept for a fractional dose schedule under conditions of natural exposure. The study will be conducted in children 5-17 months old at first vaccination living in areas of mid to high malaria transmission, in line with the age group recommended by the World Health Organization. Results from study will be critical in informing future possibilities for the development of vaccine-based strategies which, in combination with other interventions, may contribute to the malaria elimination agenda.
	CROWN CORONATION	Phase III	Covid-19	Measles Rubella Vaccine     Matching Placebo 3 AstraZeneca     vaccinel Vaccine	7th September 2020	Prof. Kwadwo Koram	Ga East Municipal HospitalKorle-Bu Teaching HospitalUGMCEffla-Nkwanta HospitalEentecost Treatment Center	Each country serves as its own sponsor but will receive funding from the Covid 19 Therapeutics Accelerator and Gates	Study ended Final report yet to be submitted 8 Months .	The purpose of this study is to determine that MR vaccine increases the likelihood of making the specific AstraZeneca COVID-19 vaccine more effective in people with prior exposure to the MR vaccine. This study has two different groups: one group will receive the active MR vaccine and one will receive a placebo. Thirty and sixty days later, participants in each group will receive the AstraZeneca COVID-19 vaccine.
	DOLF_IDA ONCHO SAFETY GHANA	Phase II	Onchocerciasis	1.Diethylcarbamazine Citrate I. P 100mg 2.lvermectin (Stromectol® 3mg) 3.Albendazole (Zentel™ 400mg) / Allopathic druss	22nd February 2019	Dr. Nicholas Oboku	University of Health and Allied Sciences	Washington University School of Medicine	Study ended Final report submitted 24 Months	Programs for control of onchocerciasis through community directed treatment with ivermectin (IVM) as a form of Mass Drug Administration (MDA) have been in place for almost 30 years. IVM is effective for clearing Mf and it temporarily sterilizes adult female worms, but it is not a microfilaricide and does not kill adult worms. For that reason, MDA with IVM must be repeated for the reproductive life of the adult worms, which is 10-15 years. Thus, there is a widely recognized need for new, safe, short-course treatment drug(s) that can kill or permanently sterilize adult worms.  This study aims to provide preliminary data on the safety of ivermectin + diethhy/carbamazine + abbendazol (IDA) treatment in persons with onchocerciasis when administered after pre-treatment with IVM to clear or greatly reduce microfilariae from the skin and yess. Widespread use of IDA following IVM pretreatment (I/IDA) has the potential to greatly accelerate elimination of LF in African countries that are covered microfilariae.
- 30	SMAART	Phase II	Stroke	1.POLYCAP 2.USUAL CARE		Dr. Fred Stephen Sarfo	Komfo Anokye Teaching Hospital	Kwame Nkrumah University of Science and Technology	Study ended Final report submitted 19 months	There has been unprecedented rise in the prevalence of stroke in sub-Saharan Africa (SSA), which when compared to stroke proffles in high-income countries (HIC) is characterized by a younger age of onset, higher case fatality rates, and more severe disability among survivors. Stroke survivors in SSA are especially at high risk for recurrent vascular events or death due to several factors including uncoordinated health systems, undiagnosed and under-controlled vascular risk factors, and lack of care affordability. Fixed-dose combination pills, known as "polypills", containing Aspirin, a statin and blood pressure (BP) lowering medication(s) may improve medication adherence and consequently reduce vascular risk as a cost-effective intervention among high risk patients including stroke survivors. This trial is to assess whether a polypill containing fixed doses of 3 antihypertensives, a statin and antiplatelet therapy taken once daily orally would result in carolid infinal thickness regression, improved adherence, and tolerability compared with usual care' group on separate individual secondary preventive medications among Ghanaian first time stroke survivors (male of refinele above the age of 18 years).
70	LEDoxy	Phase II	Lymphatic Filariasis	1.Doxycycline (Remycin®100mg 2.Placebo 3.Standard MDA Treatment/ Aliopathic drug	12th July, 2017	Prof. Alexander Yaw Debrah	Kumasi Centre for Collaborative Research (KCCR), Kwame Nkrumah University of Science and Technology (KNUST)     War Memorial Hospital, Navrongo	Kumasi Center For Collaborative Research (KCCR)	Study ended Final report submitted 40 months	The previously demonstrated effect of doxycycline in reversing or stopping the progression of lymphedema of patients with stage 1-3, irrespective of their filarial infections being active or not, provides an opportunity to include the drug as a new tool inlymphatic filariasis (LF) morbidity management programs. However, before recommendations can be made regarding the frequency of its usage or alternate dosing patterns more trials need to be conducted. This multi-national trial is to show efficacy of a lower dosage of doxycycline and to confirm finding in patients with stages 1-3 ymphedema irrespective of active LF infection as well as in people with higher grades of lymphedema.  The purpose of the study is to establish that Doxycycline can improve filarial lymphedema in healthy adolescents or adults (14 – 65 years)

:	FALCON	Phase III	Surgery	1.ChloraPrep™ stick 2.Videne® Antiseptic Solution 3.Triclosan Coated PDS and/or Vicryl sutures 4.Non-triclosan coated PDS and/or Vicryl sutures/ Medical device	10th April, 2019	Т	Tamale Teaching Hospital	The University of Birmingham	Study ended Final report submitted 24 Months	Improving surgical outcomes is a global health priority. Recent World Health Organisation (WHO) guidelines made 29 recommendations for intraoperative and postoperative measures to prevent SSI, including global perspectives relevant to LMICs., none of the evidence for the recommendations used was derived from resource limited settings, leading to uncertainty about implementation of measures in these settings. A randomised trial that has the potential to evaluate multiple interventions has particular value in this setting, and can establish a high quality evidence base that will inform guidance, and influence revisions to the WHO Surgical Safety Checklist This study assesses whether either (1) 2% alcoholic chlorhexidine versus 10% povidone-lodine for skin preparation, or (2) triclosan-coated suture versus non-coated suture for fascial closure, can reduce surgical site infection at 30-days post-surgery for each of (1) clean-contaminated and (2) contaminated/dirty surgery  The purpose of this trial is to evaluate the:
	KNC 19 (NIBIMA)	Phase IIb	Covid-19	Nibima 2.WHO standard treatment for COVID-19/ Herbal drug	11th September 2020	Prof. Ellis Owusu-Dabo		KNUST Office of Grants and Research	Study ended Final report submitted From 3 months to 7 months	-Efficacy of Nibima in reducing >50% Covid-19 viral load per patient within 14 days of therapy.  Evaluate the efficacy of Nibima in increasing the anti-inflammatory and interferon alpha/beta profiles of >50% of the Covid-19 patients within 14 days.
;	MULTIMAL 73	Phase II	Malaria	Artesunate Pyronaridine (Pyramax 2.Atovaquone Proguani (Maliarone) 3.Clindamyci 4.Eosdidomysn5.Artesunate / Allopathic drug	27th July 2020	P/(s) Dr. Oumou Maiga (KCCR)	St. Francis Xavier Hospital Assin Fosu, Ghana. Gabon	Department of Tropical Medicine, Bernhard Nocht Institute for Tropical Medicine (BNITM)	Study ended Final report submitted 7 months	The main objective of the project is to investigate two combinations of drugs already used in the market or in late-stage clinical development but not yet tested in the presently proposed combination. These are Artesunate-Pyronardin-Aloxaquone-Proguanti (APAP) and Artesunate-FosmidomycinClindamycin (AFC).  The drug combinations will be investigated in a randomized controlledthree-group clinical phase il study.  This study will aim to describe:  The pharmacokinetics of the investigated drugs when administered in combination therapy.  PCR corrected antimalarial efficacy over a 42 day follow up period  Safety and tolerability.
	74 STAR TRIAL	Phase IV	Anaesthesia	1.Paracetamol 2.Morphine/Allopathic drug	7th May 2021	Dr. Frank Enoch Gyamfi		Dr. Frank Enoch Gyamfi	Study ended Final report submittee 10 months	To compare the efficacy of intramuscular (i.m) morphine as unimodal analgesic with bimodal administration of i.m. morphine and i.v. paracetamol in managing postoperative pain in emergency abdominal surgery.  To assess the response of patients to i.m. morphine in pain management after emergency abdominal surgery.  To assess the response of patients to a combination of i.v. paracetamol and i.m. morphine in managing pain after emergency abdominal surgery.  To determine the administered analgesic and length of hospital stay.  To determine the association between administered analgesic and postoperative complications.
	DIABETIC FOOT 75 SELF CARE	Feasibility testing	Diabetes	1.Foot Selfcare Training and Education Plus usual care 2. Usual care/Training	28th October 2021	Dr.Joseph N. Suglo	Diabetes Clinic, Komfo Anokye Teaching Hospital (KATH) – Ghana	King's College London (KCL)	Study ended Final report in E3 format submitted, 7 months	The primary aim of this research is to evaluate the feasibility of conducting a randomised controlled trial to investigate the effectiveness of a hands-on skills training and education on foot self-care programme for persons with diabetes and their family caregivers in Ghana. The research question is 'can the provision of a family-oriented foot self-care skills training and education intervention improve foot care behaviour, foot care self-efficacy, knowledge of diabetic foot and diabetes distress among persons with diabetes and their caregivers in Ghana?'
;	CHEETAH 76	Pilot	Surgery	Sterile Gloves     Sterile Surgical Instrument/Medical device	1st June 2020	Professor Stephen Tabiri	Holy Family Hospital – Berekum     Holy Family Hospital – Techiman	Birmingham Clinical Trials Unit, University of Birmingham	Study ended Final report submitted. 24 Months	To purpose of this study is to assess whether the practice of using separate, sterile gloves and instruments to close wounds at the end of surgery can reduce surgical site infection at 30-days post-surgery for patients undergoing clean-contaminated, contaminated or dirty abdominal surgery, compared to current routine hospital practice.
	KAE609	Phase II	Malaria	1.KAE609 2.COARTEM TABLETS /Allopathic drug 1.Smal Quantity Lipid-based Nutrient	8th August 2017	Dr. Abraham Rexford Oduro	Navrongo Health Center	Novartis Pharma AG, Switzerland		KAE609 will be evaluated primarily for hepatic safety of single and multiple doses in sequential cohorts with increasing doses. This study aims to determine the maximum safe dose of the investigational drug KAE609 in Adult patients with acute, uncomplicated Plasmodium falciparum malaria infection
	Saving Brains Navrongo 78	Phase I	Malnutrition	Supplement for Pregnant and Lactating mothers (SQLNS P&L) 2.1 Enhanced Small Quantity Lipid-based Nutrient Supplement for Pregnant and Lactating mothers (eSQLNS P&L) 3.SQLNS for Infants 4.eSQLNS 5.SQLNS nut 6.Omega 3 flaty acids 7. Corn oil Food supplements 11. Small Quantity Lipid-based Nutrient	7th February 2019	Dr. Engelbert A. Nonterah	Navrongo Health Research Centre	Nutriset, SAS	Study ended; Final report yet to be submitted 6:months	Mainutrition continues to be a global problem. Globally 156 milion children less than 5 years are stunted, 50 million wasted, while simultaneously 42 million are overweight reflecting the double burden of mainutrition. Prevalence of mainutrition varies by region and country with Asia and Africa being the worst affected regions. This study is to seess the acceptability and adherence to nutrient supplementation for 6 weeks among pregnant and lactating women and 6 monh old infants post weaning
;	SAVING BRAINS KUMASI 79	Phase I	Malnutrition	Supplement for Pregnant and Lactating mothers (SQLNS P&L) 2. Enhanced Small Quantity lipid-based Nutrient Supplement for Pregnant and Lactating mothers (eSQLNS P&L) 3. SQLNS for Infants 4.6SQLNS for Infants 5. Omega 3 fatty acids/ Food supplements	1st November 2017	Prof. Jacob Plange-Rhule	Tafo Government Hospital     Suntreso Government Hospital     Kumasi South Government Hospital	KNUST/Nutriset SAS	Study ended 6months	Malutrition continues to be a global problem. Globally 156 milion children less than 5 years are stunted, 50 million wasted, while simultaneously 42 million are overweight reflecting the double burden of malnutrition. Prevalence of malnutrition varies by region and country with Asia and Africa being the worst affected regions. This study is to seess the acceptability and adherence to nutrient supplementation for 6 weeks among pregnant and lactating women and 6 monh old infants post weaning

80	ALB_IVM	Phase III	Onchocerciasis	Ivermectin, Albendazole Allopathic drug	1st April 2014	Dr. Nicholas Opoku	Onchocerciasis Chemotherapy Research Centre Government Hospital.	Case Western Reserve University School of Medicine, 10900 Euclid Ave Cleveland	Study ended; Final report submitted 38 months	To address whether IVM plus ALB given twice per year will be superior over annual treatment or IVM given biannually
81	MAL 055	Phase III	Malaria	RTS.S/AS01E/ Vaccine	1st October 2008	1. Prof. E. Tsiri Agbenyaga 2. Prof. Seth Owusu Agyei 3. Dr. Kwaku Poku Asante	Malaria Research Centre, Agogo.     Kintampo Health Research Centre	GlaxoSmithKline Biologicals	Study ended; Final report submitted 60 months	This Phase III study of GSK Biologicals candidate malaria vaccine RTS.S/AS01E has been designed to address the key safety and efficacy information required for vaccine licensure. In addition, other disease endpoints that allow the evaluation of the full public health impact and cost effectiveness of vaccine implementation are included. Coprimary objectives will investigate the efficacy against clinical disease in children from 5-17 months of age at first dose and the efficacy in infants 6-12 weeks of age who receive the vaccine in co-administration with EPI antigens
82	MMS	Phase III	Malnutrition	Multiple micronutrient supplement 2.lron + folic acid tablets/ Food supplements	2nd October 2012	Prof. Tsiri Agbenyaga	Barekuma Collaborative Community Development Project     C/O Komfo Anokye Teaching Hospital, Kumasi	Kirk Humanitarian	Study Ended; yet to submit report 48 months	
82	PRENABELT	Prilase III		1.Prenabelt™ 2. Sham prenabelt™     3.Body Position Sensor/ Medical device	21st April 2015	Dr. Jerry Coleman	Korle-Bu Teaching Hospital, Accra – Korle Bu	Global Innovations for Reproductive Health and Life, USA	Study ended; Final report submitted 7 months	The purpose of this study is to determine the effect of the PrenaBelt on birth- weight and assess the feasibility of introducing it to Ghanaian third-trimester pregnant women in their home setting via an antenatal care clinic and local health-care staff. Data from this study will be used in effect size calculations for the design of a large- scale, epidemiological study targeted at reducing LBW and SB in Ghana and globally.
	СРАР	Phase III	Infant Acute Respiratory Distress	1.DeVilbiss IntelliPAP CPAP machine (Model DV5 Series)     2. Hudson RCI nasal cannulas/ Medical device	14th May 2013	Dr. Harry Tagbor     Dr. Frank Baiden     Dr. Damien Punguyire	Mampong Government Hospital, Mampong     Kintampo Municipal Hospital, Kintampo	(GE) Foundation's Systems Improvement at	Study ended; yet to submit report in required format. 36 months	Evaluating the impact of using continuous positive airway pressure (CPAP) on mortality among children admitted into emergencies wards. an interventional trial to determine if CPAP reduces mortality in children 1 month to 5 years of age with acute respiratory distress
85	AIMS	Phase III	Transfusion- Transmitted Malaria (TTM)	Mirasol system for whole blood 2.Standard fresh whole blood/Blood product	9th July 2013	Dr. Shirley Owusu-Ofori	Komfo Anokye Teaching Hospital	Terumo BCT Europe N.V.	Study ended; Final report submitted 6 months	The objective of this study was to evaluate the efficacy of Mirasol-treated fresh whole blood (WB) to prevent transfusion-transmitted malaria (TTM) by comparing the incidence of TTM between subjects receiving Mirasol-treated fresh WB and subjects receiving standard (untreated) fresh WB.
86	MENINGOCOCCAL-A CONJUGATE VACCINE	Phase III	Meningitis	Meningococcal A Conjugate Vaccine/ Vaccine	26th June 2007	Dr. Patrick Ansah	Navrongo Health Research Centre	SIIL PATH	Study ended; Final report submitted 54 months	To compare the immunogenicity at 28 days after vaccination of range dosages - 10, 5, and 2.5 µg of the PsA-TT vaccine, when administered to infants in a two-dose schedule at 14 weeks (window 14 to 18 weeks of age) and 9 months of age (window 9 to 12 months of age) concomitantly with EPI vaccines (Groups 1A vs. 1B vs. 1C)
	NON-INVASIVE HAEM DEVICE	Phase III	Hemoglobin deficiency in Pregnant women	Pronto & pronto-7 pulse co-oximeter pulse co-oximeter 2. Hemocue 201+3.     Abx pentra 60 hematology analyzer/ Medical device	9th April 2013	Dr. Sam Newton	Kintampo Health Research Centre, Kintampo	РАТН	Study Ended Final report submitted 2 months	Aim The aim of the validation study was to evaluate the accuracy of the Pronto and Pronto 7devices in measuring Hb when compared to measuring Hb using the Hemocue and the ABX Pentra 60 hematology analyzer as the reference standard. Study Objectives: To compare Hb values as measured by the Pronto and Pronto 7noninvasive Hb devices and HemoCue 201+ machine with those obtained by a venous blood draw using an ABX Pentra 60 hematology analyzer among pregnant women attending ANC clinic in Ghana.
	ROTARIX	Phase III		Rotarix™/ Vaccine	6th February 2012	Prof. George Armah	Navrongo Health Research Centre	PATH	Study Ended 7 months Final Report submited	To show the superiority of live, oral Rotarix vaccine administered at 6, 10, and 14 weeks of age versus live, oral Rotarix vaccine administered at 6 and 10 weeks of age in terms of serum rotavirus immunoglobulin A (IgA) seroconversion as the marker of vaccine-induced immunogenicity
	ARTIMIST	Phase III	Malaria	ArTiMist/ Allopathic drug	22nd October 2010	Dr. Patrick Ansah	Navrongo Health Research Centre	ProtoPharma Limited	Study Ended Final report submitted 5 months	The primary objective of this study was to demonstrate the superiority of ArTiMist™ over intravenous (iv) quinine in establishing parasite success (reduction of parasite counts by ≥ 90% within ≥ 4 hours) in children with severe or complicated falciparum malaria, or children with uncomplicated malaria with gastrointestinal complications.
90		Phase III	Human Papilom Virus (HPV)	Gardasil/ Vaccine	1st November 2010		Navrongo Health Research Centre	Merck, Sharp and Dohme Corporation	Study Ended Final report submitted 20 months	To estimate the percentage of subjects who seroconvert to each of HPV 6, 11, 16, and 18 at Month 7 (4 weeks Postdose 3).  To evaluate the safety and tolerability of CARDASIL in females 9 to 26 years of age in SubSaharan Africa.  Secondary: To estimate Month 7 anti-HPV 6, 11, 16, and 18 geometric mean titlers (GMTs) in vaccinated subjects.

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01	SMAC	Phase III	Malaria	Intravenous Artesunate 2.     Intramuscular Artesunate/ Allopathic	1st January 2013	Prof. Tsiri Agbenyega	Komfo Anokye Teaching Hospital,	University Medical Centre Tubingen	Study Ended 15 months	
31	SWAC	riiase iii	ivididid	Intrantuscular Artesuriate/ Aliopatric	ist January 2013	Prof. Islii Agueriyega	Kuillasi	Tubligen	13 monus	
92	OXYTOCIN	III	Postpartum Hemorrhage (PPH)	1.Oxytocin in uniject™ 10 iu/ Hormone	12th May 2010	Dr. Sam Newton	Kintampo Health Research Centre	PATH	Study Ended Final report submitted 12 months	To determine the effect of prophylactic administration of oxytocin in uniject on postpartum haemorrhage at home births in the Kintampo north and south districts of Ghana
	AMARYL M		Type 2						Study Ended	To determine the clinical Efficacy and Safety of Amanyl M in Patients with Type 2 Diabetes Who are Inadequately Treated by Either Glimepride or Metformin Monotherapy or Who are Afready Treated with Free Combination of Glimepride and
93	3	IV	Diabetes	Amaryl m oral tablets/ Allopathic	16th October 2009	Dr. Frank Umeh	Korle-Bu Teaching Hospital	Sanofi Aventis	6 months	Metformin in African Countries
	MOXIDECTIN- IVERMECTIN						Onchocerciasis Chemotherapy Research Centre Government	Wyeth Research Division of Wyeth Pharmaceuticals Inc.     Product Development and Evaluation unit TDR	Study Ended Report submitted	To determine the Safety, Tolerability, and Efficacy of Orally Administered Moxidectin
94		III	Onchocerciasis	1. Moxidectin 2. Ivermectin/Allopathic	1st February 2004	Dr. Nicholas Opoku	Hospital.		25 months + (12 months ext.)	in Subjects with Onchocerca vovulus
								Wyeth Research Division of Wyeth Pharmaceuticals Inc.		
95	MOXIDECTIN	Phase II	Onchocerciasis	Moxidectin 2mg Tablets/Allopathic	1st February 2004	Dr. Kwabla Awadzi	Onchocerciasis Chemotherapy Research Centre Government Hospita	2. Product Development and Evaluation unit TDR	Study Ended Ended 60 months	
96	EBA	Phase I	Malaria	(EBA-175 RII-NG) malaria vaccine/ Vaccine	1st March 2009	Prof. Kwadwo Ansah Koram	Noguchi Momorial Institute of Medical Research	Division of Microbiology and Infectious Diseases (DMID) National Institute of Allergy and Infectious Diseases (NIAID)	Study Ended Final report submitted 18 months	To determine the Immunogenicity of EBA-175 RII-NG Malaria Vaccine Administered Intramuscularly in Semi-Immune Adults
0.7	IPT & SP	Dhara III	Malaria in Pregnant		4.4440000	S. Abarbara Hadasara	Health Facilities in the Kassena Nankana, Navrongo Health Research	London School of Hygiene and Tropical	Study Ended	to compare the intermittent preventive treatment of sulfadoxine-pyrimethamine with
97		Phase III	women	Sulfadoxine-pyrimethamine/Allopathic	1st May 2008	Dr. Abraham Hodgson	Centre	Medicine	32 months	intermittent screening and treatment of malaria in pregnancy
98	IRON FORTIFICATION III		Malaria	1.Sprinkles vitamine 2.mineral food supplement/ Food supplements	1st July 2009	Prof. Seth Owusu Agyei	Kintampo Health Research Centre	National Institutes of Health	Study Ended 12 months	To determine the seasonal impact of iron fortification on malaria incidence in Ghanaian children
	ROTASHIELD		Rotavirus			Prof. George E. Armah     Prof. Fred N. Binka     Dr. Abraham Hodgson	War Memorial Hospital, Navrongo     Bongo Hospital	International Medica	Study Ended	To determine the efficacy, immunogenicity, and safety of two single doses of RRV TV
99		III	Gastroenteritis	RRV-TV Vaccine (rotashield)/ Vaccine	1st August 2009			Foundation	16 months	in neonates / infants
100	AZITHROMYCIN PLUS CHLOROQUINE PHOSPHATE	Ш	Malaria	Azithromycin 2. Chloroquine Phosphate     Artemether- Lumefatrine/Allopathic	1st October 2007	Dr. Patrick Ansah	Navrongo Health Research Centre	Pfizer Laboratories Incorporated, Pfizer Global Research and Development.	Study Ended Final report submitted 8 months	To compare azithromycin plus chloroquine phosphate with artemether-lumefantrine for the treatment of uncomplicated plasmodium falciparum malaria in children in Africa
101	CRASH-2	I	Trauma patient with or at risk of hemorrhage	1.Tranexamic acid 2. Placebo/	1st August 2007	Prof. J. C. B. Dakubo	Korle-Bu Teaching Hospital	London School of Hygiene & Tropical Medicine	Study Ended, Lancet publication submitted 24 months	To determine the effects of anti-fibrinolytic treatment on death and transfusion requirement among trauma patients with or at risk of significant haemorrhage.

	F	PYRONARIDINE									
	A	ARTESUNATE VRS COARTEM			1.Pyronaridine Artesunate Tablet (PYRAMAX)				Medicines For		To Compare the Safety and Efficacy Of Fixed Dose Formulation Of Oral Pyronaridine
	102		ш	Malaria	2.Artemether-Lumefantrine(COARTEM)/ Allopathic	1st March 2007	Dr. G. Bedu-Adoo	Komfo Anokye Teaching Hospital	Malaria Venture, Switzerland	Study Ended 3 months	Artesunate Tablet with Coartem In Children And Adult Patients With Acute Uncomplicated Plasmodium Falciparium Malaria
	102			ivididild	Ліюранію	TSUMAICH 2007	Dr. G. Bedu-Adoo	Rollio Allokye Teaching Hospital	Swizeriariu	3 monuis	Oncomplicated Flashfodum Falcipanum Malana
		MAL 050									
		WAL 050							GlaxoSmithKline	Study Ended	
1	103		<u>                                     </u>	Malaria	RTSS, AS10E Vaccine/Vaccine		Prof. Seth Owusu Adjei	Kintampo Health Research Centre	R&D	17 months	
									Division of Microbiology		
									and Infectious Diseases		
									(DMID)		
									National Institute of Allergy and		
	F	PFCSP_MVACS_MA _ARIA							Infectious Diseases		
		LAKIA							(NIAID)	Study Ended	
1	104		1	Malaria	PfCSP DNA VACCINE (VCL-2510)/Vaccine	1st August 2005	Prof. Kwadwo A Koram	Tetteh Quarshie Memorial Hospital		18 months	
	F	ROTATEQ							1. Merck & Co.	Study Ended Final report	
1	105		Ш	Gastroenteritis	Rotateq/Vaccine	1st September 2007	Prof. George E. Armah	Navrongo Health Research Centre	2. PATH	published in Lancet 18 months	
	N	MEFLOQCHLOAZITH								Study Ended Final report	
1	106		Ш	Malaria	Mefloquine 2. Chloroquine 3.     Azythromycin/Allopathic	4th August 2004	Dr. Abraham Hodgson	Navrongo Health Research Centre	Pfizer Inc.	submitted 12 months	
						<b>y</b>	,				
		MAL 047					Prof. Seth Owusu Adjei,				
	107	VIAL 047		Malaria	1.RTS,S/AS02D 2.RTS,S/AS01E/Vaccine		Dr. Kwaku Poku Asante	Kintampo Health Research Centre	GlaxoSmithKline R&D	Study Ended 19 months	
,	107		<u> </u>	Ivialaria	1.R15,5/A502D 2.R15,5/A501E/Vaccine			Kintampo nealin Research Centre	RaD	19 MONUIS	
	c	CDA					Prof. Seth Owusu Agyei				
	108		ш	Malaria	1.Chorproguanil-Dapsone-Artesunate (CDA)     2.Artemether-Lumefantrine/Allopathic	19th July 2006	Dr. Kwaku Poku Asante	Kintampo Health Research Centre	GlaxoSmithKline R & D	Study Ended 12 months	
	100			Walana	2.Artemetici-Eurocantino/Aiopatilo	1301 July 2000		Trintampo ricalar rescalon ocnite	Kub	12 mondis	
	_	CDA2									
		JUNE		Malaria	1. Chorproguanil-Dapsone-Artesunate (CDA)	27 luna 2006	Deaf Tairi Ashanyas	Department of Physiology, School of Medical Sciences, KNUST	GlaxoSmithKline	Study Ended	
1	109		III .	ivialaria	z.Artemetrer-Lumerantrine/aliopatriic	Z1,Juile 2006	Prof. TSITI Agbenyega	INIEUICAI SCIENCES, KINUS I	K & D	12 months	
									United States		
									Agency for International		
									Development		
									Through The		
		JOVA OII					Durk Durkligt		Collaborative		
		NOVASIL					Prof. David Ofori Agyei Dr. Nii- Ayi Ankrah	Ejura Sekyedumasi Disrict, Ashanti	Support	Study Ended	
1	110		II		NovaSIL			Region	Program	9 months	
	Т	TENOFOVIR							Family Health	Study Ended	
1	111		II	HIV	Tenofovir Disoproxyl Fumarate (TDF)/Vaccine	1st February 2004	Dr. Edith Clarke	Ghana Health Service	International	20 months	
	110	NOVASIL FENOFOVIR	III	Malaria	2. Artemether-Lumefantrine/allopathic			Ejura Sekyedumasi Disrict, Ashanti Region	R & D  United States Agency for International Development (USAID) Through The Peanut Collaborative Research Support Program	Study Ended 9 months	

							Dr. William Ampofo	Noguchi Memorial Institution for Medical Research.			
	SAV	VVY					Dr. Baafuor Kofi Opoku	iwedical Research.			
	40				SAVVY (Microbicide)	1st February 2004		Komfo Anokye Teaching Hospital.	Family Health International	Study Ended 32 months	
1	12		<u> </u>		SAVVY (MICRODICIDE)	1st February 2004			international	32 months	
	MAI	L 063							Malaria	Study Ended Final report	
1	13		Ш	Malaria	RTS,S/AS01E/ Vaccine	15th April 2011	Prof. F. Tsiri Agbenyaga		Research Centre, Agogo	submitted 52 months	
			<del></del>				gy-g-				
					Eurartesim oral tablets     Farmanguinhos artesunate+mefloquine			Ejisu Government Hospital, Ejisu	Prince Leopold		
	PRE	EGACT			fixed combination oral tablets		1.Dr. Harry Tagbor	Juaben Government Hospital,	Institute of		
1	14		Ш		Coarsucam oral tablets/ Allopathic		2.Dr. Henry Opare Addo	Juaben	Tropical Medicine	Study Ended 60 months	
										Study Ended, Yet to submit final	
	ALE	BIVIM K'SI							University	report	
	15			Onchocerciasis	I. Ivermectin     2. Albendazole/Allopathic	10th Nevember 2015	Prof. Alexander Yaw Debrah	Kumasi Centre for Collaborative Research in Tropical Medicine	Hospitals Case medical Center	4 years and 2 months	
1	15		III	Offchocerciasis	1. Ivermecum 2. Albendazoie/Aliopathic	Total November 2015	Debian	Research in Tropical Medicine	medical Center		
	RIF.	FAMPIN VS ONIAZID									
	ISO	DNIAZID			1.lsoniazid 2. Rifampin/Allopathic/				Canadian Institute of	Study Ended 60 months	
1	16		III	Tuberclosis	Allopathic	2nd March 2011	Dr. Joseph Baah Obeng		Health Research		
	NO	GUCHI			Alere filariasis test strip     2.Sd bioline		Prof. Daniel A. Boakye			Study Ended Final report	
	FIL/	ARIASIS			lymphatic filariasis IgG4 3.Sd bioline oncho/lf IgG4 biplex		Dr. Nana – Kwadwo Biritwum	Noguchi Memorial Institute For Medical	World Health		Development of a plan of action for strengthening LF elimination in Ghana, and where appropriate, a plan of action for integrating LF and onchocerciasis elimination efforts,
1	17			Filariasis	4.Diethylcarbamazine patch /Allopathic	7th June 2017	Biritwum		TDR		appropriate, a plan of action for integrating LF and onchocerclasis elimination efforts, to be proposed to the GHS decision makers.
											To evaluate the safety of 1.25mg and 2mg ziv-aflibercept in Ghanaian population with retinal vascular diseases.  To determine the safety of intravitreal injections
											of ziv-affibercept at 4 and 12 weeks in a Ghanaian population.
	70.4	(AFELIDEDOEST		Dational						Study Ended Final report	To measure the visual outcome of treatment with 1.25mg and 2mg ziv-aflibercept in
	ZIV	AFFLIBERCEPT		Retinal Vascular				Retina unit, Eye Centre, Korle-Bu,			eyes with DME, nvAMD, and ME secondary to RVO at 12 weeks.  To measure the anatomic changes using SD-OCT in eyes with DME, nvAMD and ME
1	18		I .	diseases	1.Ziv-aflibercept (ZALTRAP) / Allopathic	30th January 2017	Braimah Imoro Zeba	Teaching Hospital, Korle-Bu, Accra	Same as PI		secondary to RVO at 12 weeks.
											Sickle cell disease (SCD) is a genetic, autosomal, recessive blood disorder resulting in
											altered (sickle- shaped) red-blood cells. A vaso-occlusive crisis (VOC) is a severe, acute painful episode that occurs when sickle-shaped red blood cells obstruct the
											microcirculation and restrict blood flow to an organ or tissue, resulting in ischaemia,
								Komfo Anokye Teaching Hospital,			necrosis and organ damage. There is a high unmet need for treatment options in SCD and there is a data that platelet inhibition has the potential to reduce the risk for acute
								Department of Child Health			and there is a data that platelet inhibition has the potential to reduce the risk for acute vaso-occlusions.
							Prof. Alex Osei-Akoto     Dr Patrick Ansah	Navrongo Health Research Centre     Department of Child Health, Korle			This study is to evaluate the effect (efficiency selects and teleschills)
								Department of Child Health, Korle     Bu			This study is to evaluate the effect (efficacy, safety and tolerability) of ticagrelor versus placebo in reducing the rate of vaso-occlusive crises (VOCs), which is the composite
	us	STIA3	Phase III	Sickle Cell			4.Dr Kokou Hefoume	University of Health and Allied		submitted	of painful crisis and/or acute chest syndrome (ACS), in paediatric patients (2 to 11
1	19 HE	STIAS	rnase III	Disease	1.Ticagrelor 2.Placebo/Allopathic	1st August, 2018	Amegan-Aho	Sciences	AstraZeneca AB	29 MONTHS	years and 12 to 17 years with sickle cell disease (SCD).
						9,					

				I I			I			
12	PRCR DIPSTICK	Phase II	proteinuria	1.Test-It™ Protein Creatinine Dipstick 2.Urinalysis Reagent Strips 3.Quantitative Spectrophotometric Method/Medical device	16th February, 2018	Dr. Sam Newton	Kintampo Health Research Center	Program For Appropriate Technology In Health (PATH)	Study Ended. Final Report Submitted 19 months	The lack of access to reliable tests for proteinuria measurement in all antenatal care settings, particularly at the periphery, remains a critical gap in the accurate identification of women at high risk for Pre-Eclampsia. In Low Resource Settings, a protein-only measurement via a urine dipstick is the most widely used proteinura test due in part to its low complexity and low cost. However, the clinical utility of the protein-only dipstick is limited. Test results can be unreliable, as the test cannot adjust for daily fluctuation of body hydration. This leads to protein measurements that are either too low or too high due to the level of urine dilution. More accurate tests, such as the 24-hour urine test, are available only for confirmatory testing in tertiary-level clinics due to their high cost and technical complexity.  The purpose of the study is to generate a body of evidence that will determine performance characteristics of the current Protein Creatinine dipstick test and the feasibility of its use in target Ante Natal Care settings.
12	MAL 073	Phase IIIb	Malaria	1.RTS,S/AS01E 2.MR-VAC™ 3.STAMARIL4.VITAMIN A /Vaccine	11th December 2015	1.Prof. Tsiri Agbenyega Prof. Seth Owusu Adjei	Malaria Research Center, Agogo     Kintampo Health Research Centre		Study Ended Final Report submitted 43 months 16 days	In sub-Saharan Africa, most of the Expanded Program on Immunization (EPI) vaccines are given in early infancy while measles, rubella and yellow fever (YF) vaccines are given at 9 months of age. Between the first EPI vaccines and the measles, rubela and YF vaccines, children receive Vitamin A supplementation at 6 months of age. To limit the number of ciline visus for young other and to optimize vaccine implementation a schedule (0, 1.5, 3-month) is proposed.  There are however no data of the anti-circumsportozoite protein of Plasmodium falciparum (anti-CS) immune response induced by RTS,S/ASO1E when given in co-administration with measles, rubella and YF, in a 0, 1.5, 3-month schedule starting at an older age (5-17 months). This study intends to demonstrate that anti-CS immune responses of the candidate malaria vaccine RTS,S/ASO1E is not inferior when RTS,S/ASO1E is administered at 6, 7.5 and 9 months of age with the third dose given alone or in co-administration with a YF vaccine and a combined measles and rubella vaccine  Safety hornations the starting at 6 months of age. This study will therefore provide safety information when RTS,S/ASO1E is and ministered at 6, 7.5 and 9 months of age alone or in co-administration with YF vaccine and a combined measles and rubella vaccine
12	CEPHEID XPERT HIV-	PILOT	Michaelia HIV	Xpert HIV-1 VL XC Test Assay for detecting	6th June 2019	Prof. Jacob Plange-Rhule	St. Martin De Porres Hospital Atua Government Hospital Akosombo Hospital	CEPHEID	Study Ended Final Report yet to be submitted 6 Months	The Xpert® HIV-1 Viral Load XC test is an in vitro reverse transcriptase polymerase chain reaction (RT-PCR) assay for the quantification of Human Immunodeficiency Virus type 1 (HIV-1) RNA in human plasma using the automated GeneXpert® Instrument Systems. It is intended for use as an aid in the diagnosis of HIV-1 infection, as a confirmation of HIV-1 infection, and as an aid in clinical management of patients infected with HIV-1.
12	3 GBT440-038	Phase III	Sickle Cell Disease	Voxelotor (GBT440) Alliopathic Oral	10th February 2023	Dr. Catherine Segbefia     Dr. Vivian Paintsil		Global Blood Therapeutics, Inc.	Application closed by sponsor before commencement , 24months	The objective of this OLE is to assess the safety of, and SCD related complications with, long term treatment with Vovelotor in pparticipants who have completed treatment in a GBT-spnsored voxelotor clinical study based on the following parameters a) Adverse Events (AEs), Clinical Laboratory Tests, Physical Examinations (PEs) and other clinical measures.  b) Frequency of SCD-related complications.
12	4 CIELO Trial	Phase III	Encephalitis	Satralizumab Monoclonal antibody Subcutaneous injection through thigh/abdomen	20th December 2022	Prof. Fred Stephen Sarfo	Komfo Anokye Teaching Hospital (KATH)	F-Hoffman LA Roche/ Chugai Pharma Co. LTD	Application closed by sponsor before commencement Syears Smonths	This study will evaluate the efficacy, safety, pharmacokinetics, and pharmacodynamics of satralizumab compared with placebo in each of the following cohorts:  **NMDAR autoimmune encephalitis (AIE) cohort: adults and adolescents with definite or probable NMDAR encephalitis addition, the study will assess the long-term safety and efficacy of satralizumab during an optional extension period. For efficacy analyses, each cohort will be treated as a separate population and will have independent Type I error control at a 5% significance level Specific primary and secondary objectives and corresponding endpoints for the study are outlined below.
12	5 BLMs4BU	Phase III	Buruli Ulcer	Combination of rifampicin , Clarithromycin and Amoxicillin/Clavulanate Allopathic drug	1st February 2023	Prof. Richard Odame Phillips	St. Peters Catholic Hospital Jacobu Nkawie Government Hospital	University of Zaragoza (UNIZAR) Spain	Application closed by sponsor before commencement 2 year 11 months	The aim of this study is to determine the ability of amoxicillin/clavulanate combination therapy with rifampicin plus clarithromycin to improve the cure rate of Buruli ulcer (BU) disease compared to a standard regimen of rifampicin plus clarithromycin. Primary objective The primary objective The primary objective of this clinical trial is to demonstrate the non-inferiority of 4-week coadministration of amoxicillin/clavulanate ((RAIK/CLVI)) with rifampicin-clarithromycin ((RIF/CLA's)) compared to the standard 8-week rifampicin-clarithromycin (RIF/CLA's) in cure rates at 12 months post initiation of treatment, thus reducing BU treatment from 8 to 4 weeks.

										Primary Objective
										To evaluate the safety of mepiazumab in an adult population with uncomplicated, symptomatic <i>P. falciparum</i> infection SecondaryObjective:  To evaluate the efficacy of mepiazumab as defined by 0 Early treatment failure 0 Late parasitological failure 0 Late parasitological failure 0 Loncorrected ACPR
				Ketantin (Meplazumab) Monoclonal Antibody		Dr. Patrick Odum Ansah	Navrogo Health Research Centre (NHRC)	Jiangsu Pacific Meinuoke Biopharmaceutic	Application terminated by sponsor before commencement,	To evaluate PRR  - To determine the recrudescence ) and re-infection  - To determine the time to relief of fever  - To determine the dose-response trend relationship between 3 dose levels of meplazumab by evaluation of safety, efficacy and ACPR outcomes  - To evaluate the pharmacokinetics of meplazumab in serum
126	MPZ STUDY	Phase IIa	Malaria	Intravenous infusion	5th December 2023	2. Dr. Oumou Maiga	St. Francis Xavier Hospital/KCCR	al Co., Ltd	22 months	To evaluate immunogenicity following meplazumab administration
127	GBT-2104-133	Phase III	Sickle Cell Disease	Inclacumab/ Monoclonal antibody	27 <sup>th</sup> August, 2021	Professor Alex Osei-Akoto	Komfo Anokye Teaching Hospital (KATH)	Global Blood Therapeutics, Inc.		The primary objective of this study is to evaluate the long-term safety of every 12-week dosing of inclacumab in participants with sickle cell disease (SCD) who have completed a prior inclacumab clinical trial. Additional objectives are to evaluate the incidence of vaso-occlusive crises (VOCs), hospitalizations, missed work/school days, red blood cell (RBC) transfusions, and quality of life (QoL) with long-term use of inclacumab.
128	GBT-2104-132	Phase III	Sickle Cell Disease	Inclacumab 2.Placebo/ Monoclonal antibody	5th July, 2021	Professor Alex Osei-Akoto	Komfo Anokye Teaching Hospital (KATH)	Global Blood Therapeutics, Inc.	Study terminated by sponsor before commencement	The primary objective of this study is to evaluate the safety and efficacy of a single dose of inclacumab compared to placebo to reduce the incidence of re admission to a healthcare facility for a vaso-occlusive crisis (VCO: plater an admission for an index VOC in participants with sickle cell disease (SCD). Additional objectives of the study are to evaluate the pharmacokinetics (PK) and pharmacodynamics (PD) of inclacumab, the presence of anti-drug antibodies (ADAs), and changes in quality of life (QOL).
										The primary objective of this study is to evaluate the safety and efficacy of treatment every 12 weeks with inclacumab to reduce the incidence of VOCs in participants with SCD.
			Sickle Cell				Komfo Anokye Teaching Hospital	Global Blood Therapeutics,	before commencement	Additional objectives of the study are to evaluate the pharmacokinetics (PK) and pharmacodynamics (PD) of inclacumab, the presence of anti-drug antibodies (ADAs),
129	GBT 2104-131	Phase III	Disease	Inclacumab 2.Placebo/ Monoclonal antibody	5th July, 2021	Professor Alex Osei-Akoto	(KATH)	Inc.	2 years	and changes in quality of life (QOL).
130	INNOVATE	Phase III/II	Covid-19	1. lnn0-4800 2. Placebo/Vaccine		Susan Adu-Amankwah	Noguchi Memorial Institute for Medical Research	Inovio Pharmaceuticals , Inc	Study Closed/withdrawn by Sponsor 24 months	Evaluate the cellular and humoral immune response to INO-4800 administered by ID injection followed immediately by electroporation EP 2. Evaluate the efficacy of INO-4800 in the prevention of COVID-19 disease in subjects who are SARS-CoV-2 negative at baseline
131	LIVZON	Phase III	Covid-19	1.SARS-CoV-2 fusion protein vaccine (code: V-0)     2. Placebo/Vaccine	2nd August 2021	1.Dr Seyram Kaali 2.Dr. Nana Akosua Ansah	Mintampo Health Research Centre     Navrongo Health Research Centre	Livzon Mabpharm Inc. Institution Pharmaceutical company	Study Closed by Sponsor before commencement. No recruitment	Efficacy:  To evaluate the efficacy of the recombinant SARS-CoV-2 fusion protein vaccine (V-01) for the prevention of symptomatic RT PCR positive COVID-19 (mild or above severity) starting from at least 14 days (215 days) after full-course immunization (completing all vaccinations) Safety:  To evaluate the incidence of adverse events (AEs) of recombinant SARS-CoV-2 fusion protein vaccine (V-01) from the first vaccination to 28 days after full-course immunization
131	LIVZOIV	i ilase ili	GOVIG-13	1 BOODO/VECCITO	Zila August 2021	Z.DI. IValia Akosua Alisali	2.Mawongo Ficalar Nescaron Ochac	company	monuis	vaccination to 20 days after idir-course infindingation
132	COVID 19 INTRANASAL SPRAY	Phase III	Covid-19	I.Influenza Virus Vector COVID-19 Vaccine for Intranasal Spray     2. Placebo/Naccine	19th October 2021	Dr. Seyram Kaali	KHRC 2. NHRC     KCCR 4. Dodowa     Health Research Center     Schana Infectious Disease Center     KBTH	Beijing Wantai Biological Pharmacy Enterprise Co, Ltd		To evaluate the protective efficacy of DelNS1-2019-nCoV-RBD-OPT1 for preventing virologically confirmed (RT-PCR positive) symptomatic COVID-19.  2. To evaluate the safety of DelNS1-2019-ROOV-RBD OPT1.
			Sickle Cell				Ghana Institute of Clinical Genetics Korlebu Sickle cell office Directorate		commenced 21	The purpose of this study is to explore the effect of P-selectin inhibition with orizanlizumab on renal function in SCD patients with CKD who are receiving standard of care for SCD-related CKD, have Grade A2-A3 albuminuria and Stage 1-3a CKD,
133	STEADFAST	Phase II	Disease	CRIZANLIZUMAB/ Monoclonal antibody	30th October, 2020	Dr. Yvonne Dei Adomako	Child(KATH)	Novartis Pharma		and are at risk for rapid decline in their eGFR.
	ESM UBT		Postpartum					Bill and Melinda Gates Foundation,	Study not conducted; Funds from Sponsor withdrawn before initiation	
134	*		Hemorrhage	Uterine balloon tamponade/Medical device	17th February, 2014	Dr. Ivy Frances Osei	Field Work	USA	8months	
						Dr. Josephine C. Ocran		Sanofi-Aventis	Study Closed by Sponsor. No	
135	FERROQUINE	II	Malaria	Ferroquine 2.Amodiaquine 3.     Artesunate/Allopathic	4th January 2008	Prof. Kwadwo Ansah Koram	Noguchi Memorial Institute of Medical Research	Recherché And Development	recruitment was done. 13Conths	

136	HOPE SCD	Ш	Sickle Cell Disease	GBT440 300mg /Allopathic	1st May 2017	1.Dr. Yvonne Dei Adomakoh 2.Dr. Vivian Paintsil	Center for Clinical Genetics, Korle- Bu Teaching Hospital     Paediatric Sickle cell clinic, Komfo Anokye Teaching Hospital	Global Blood Therapeutics Inc. 400 East Jamie Court, Suite 101 South San Francisco, CA 94080,USA	Group 1 and 2 under current protocol completed (none recruited in Ghana); yet to start Main Population Study (Group 3)	The primary objective is to assess the efficacy of GBT440 in adolescents and adults with SCD as measured by improvement in anemia
497	NOSA STUDY	Phase III	Malana		Oth Neurophy 2023			Doubles	Application Consider Assessed	Primary The primary objective is to evaluate the clinical efficacy, as assessed by time to lesion(s) resolution, of IP + Standard of Care (SOC) compared to placebe + SOC for subjects with monkeypox.  To evaluate the safety and efficacy, as assessed by mortality, hospitalization, complications, and duration of symptoms of IP + SOC compared to placebe + SOC in subjects with mpox. The safety objectives are to evaluate the safety and tolerability in terms of AEs and SAEs occurrence frequencies and treatment discontinuation of 1/IP + SOC compared to placebe + SOC in subjects with non-severe mpox diseases 2/IP + SOC in subjects with non-severe mpox diseases 2/IP + SOC in subjects with severe complications and/or severe immune suppression and/or pregnancy/breastfeeding.
137	MOSA STUDY		Monkey pox	Tecovirimat	9th November, 2023			Panther	Application Pending Approval	
138	GBT021601-021	Phase II/III	Sickle Cell Disease	Osivelotor (PF-07940367/GBT021601) Allopathic drug Oral	2nd May 2024	1. Prof. Alhassan Abdul- Mumin 2. Dr. Kokou Amegan-Aho	Trafalgar Campus, Ho-Denu Road, Ho, Volta Region, Ghana     Salaga Road, Tamale, Ghana.	Global Blood Therapeutics, Inc. a wholly owned subsidiary of Pfizer	Application Withdrawn before approval, 42 Months	Primary:  To assess the effects of osivelotor in adult participants with SCD as measured by change in hemoglobin (Hb).  Part B: To assess the effects of osivelotor (adults: 150 mg QD dose) compared to placebo in adult and adolescent participants with SCD as measured by Hb response and rate of vasoocclusive crisis (VOC) events.  Part C: To assess the PK of single and MD of osivelotor in pediatric participants with SCD
139	MITAPIVAT	Phase IVIII	Sickle Cell Disease	Mitapivat Allopathic Drug Oral	24th November 2023	Dr. Eunice Agyeman Ahmed	Komfo Anokye Teaching Hospital (KATH)	Agios Pharmaceuticals , Inc	Application Withdrawn before approval,	Primary Objectives To determine the recommended Phase 3 dose of mitapivat by evaluating the effect of 2 dose levels of mitapivat versus placebo on: Anemia in subjects with sickle cell disease (SCD) Safety Secondary Objectives To evaluate the effect of 2 doses of mitapivat versus placebo on: Anemia Markers of hemolysis and erythropoiesis Patient-reported fatigue Sickle cell pain crises (SCPCs) To evaluate the pharmacokinette and
140	PROFUSA		sepsis from pulmonary or wound sources	Lumee Oxygen Sensor  Medical Device  Subcutaneous injection	12th July 2024	Dr. George Oduro	Komfo Anokye Teaching Hospital (KATH)	Henry M. Jackson Foundation for the Advancement of Military Medicine	Application Withdrawn before approval,	Primary Objective: Compare subdermal tissue oxygen concentrations in core and peripheral body sites measured via the oxygen biosensor platform with systemic blood oxygen levels in participants with suspected sepsis from pulmonary or wound sources  Secondary Objective  - Evaluate variations in tissue oxygen concentration dynamics using the oxygen biosensor platform in patients with differing sources of sepsis  - To evaluate the safety and tolerability of the biosensor technology
141	PEARL STUDY	Phase III	Respiratory Syncitial Virus	RSVI Vaccine	16th October 2023	Dr Seyram Kaali     Dr. Kokou Amegan-Aho     3. Dr. Alberta Amu     4. Dr. John Amuasi     5. Dr. Patrick Ansah     6. Prof. Tsair Agbenyeg	1. KHRC 2. UHAS 3. DHRC 4. KCCR 5. NHRC 6. Malaria Research Centre Agogo.	Sanofi Pasteur Inc	Application Withdrawn, 2 years 11 months	Efficacy  demonstrate the clinical efficacy of RSVt vaccine for the prevention of RT-PCR confirmed RSV LRTD after 2 doses, over RSV Season 1 2. To demonstrate the clinical efficacy of RSV tvaccine for the prevention of RT PCR confirmed RSV LRTD after 2 doses over RSV Season 1 3. To demonstrate the clinical efficacy of RSV tvaccine for the prevention of RT-PCR confirmed RSV associated with the occurrence of LRTD, leading to hospitalization after 2 doses over RSV Season 1 Safety To describe the safety profile of the RSVt vaccine.  Immunogenicity To describe the RSV A and B serum-neutralizing and RSV serum anti-F IgA and IgG antibody responses to the study intervention

142	ABDOV COVID-19 TRIAL	Phase III	Covid-19	SCTV01E (A COVID-19 Alpha/Beta/Delta/Omicron Variants S-Trimer Vaccine) Vaccine		1. Dr. Alberta Amu 2. Dr. Patrick Ansah 3. Dr. John Amusi 4. Dr. Kwaku Poku Asante	Dodowa Health Research Centre     Navrongo Health Research Centre     Kumasi Center for Collaborative Research (KCCR)     Kintampo Health Research Centre	Sinocelltech Ltd	Application Withdrawn, 19 Months	Stage 1 immunization evaluate the protective efficacy of SCTV01E against symptomatic COVID-19 occurring from 14 days after the 2nd dose in population previously unvaccinated with COVID-19 vaccine.  To evaluate the protective efficacy of SCTV01E, severe and above COVID-19, nospitalization due to COVID-19, and death due to COVID-19 occurring from 14 days.  To evaluate the protective efficacy of stage 1 immunization against different SARS-COV-2 variants.  Stage 2 immunization To evaluate the protective efficacy of SCTV01E against symptomatic COVID-19 occurring from 7 days after the 3rd dose in population previously unvaccinated with COVID-19 vaccine To evaluate the protective efficacy of SCTV01E, severe and above COVID-19, of SCTV01E against symptomatic COVID-19 occurring from 7 days after the 3rd dose, respectively, in population previously unvaccinated with COVID-19 vaccine To evaluate the protective efficacy of stage 2 immunization against different SARS-COV2 variants.
143	VERO CELL COVID 19 TRIAL	Phase III	Covid-19	Inactivated (Vero Cell)/Vaccine	10th February 2022	Dr Alberta Amu     Dr. Patrick Ansah	Dodowa Health Research Center     Navrongo Health Research Center	Institute of Medical Biology Chinese Academy of Medical Sciences	Application Withdrawn, 18 Months	1.To evaluate the efficacy of SARS-CoV-2 Vaccine (nactivated (Vero Cell) against symptomatic and laboratory-confirmed (RT PCR method) COVID-19 cases 2.To evaluate the solicited AEs within 7 days after each dose. 3.To evaluate the efficacy of SARS-CoV-2 Vaccine, (nactivated (Vero Cell) after at least one dose of immunization. 4. To evaluate the efficacy of SARS-CoV-2 Vaccine, inactivated (Vero Cell) against symptomatic and laboratory-confirmed (RT-PCR method) severe COVID-19 cases. 5. To evaluate the efficacy of SARS-CoV-2 Vaccine, inactivated (Vero Cell) for symptomatic and laboratory confirmed (RT-PCR method) COVID-19 cases caused by different SARS CoV-2 variantee.
144	MEBENDAZOLE	IV	Hookworm infection	Menbendazole/Allopathic			Kintampo Health Research Centre	Program For Appropriate Technology In Health (PATH)	Application Withdrawn	global health problems, thought to parasitize some 2 billion people worldwide.[] The most recent estimates suggest that between 600 and 800 million people are infected with one or several of the common soli-transmitted helminitis (STHs), which are Ascaris lumbricoides, Trichuris trichiura, and hookworm.] Infection prevalence, incidence, and disease burden are particularly high in tropical and subtropical areas that are already burdened with poor living conditions, over-population, and inadequate sanitation, including some areas of sub-Saharan Africa, Asia, and Latin America [1, 1]
145	EBOLA Z	11	Ebola	chimpanzee adenovirus Type 3 – vectored Ebola Zaire vaccine (ChAd3-EBO-Z)/Vaccine		1.Dr. Kwaku Poku Asante 2.Prof. Kwadwo A Koram	Kintampo Health Research Centre	GlaxoSmithKline Biologicals	Application withdrawn	
146	EBOLA Z (Paediatric)	11	Ebola	chimpanzee adenovirus Type 3 – vectored Ebola Zaire vaccine (ChAd3-EBO-Z)/Vaccine	21st August 2015	Dr. Kwaku Poku Asante	OCRC, Hohoe	Glaxosmithkline Biologicals, Rue De L'insitut, 89 – 1330 Rixensart, Belgium	Application withdrawn N/A	

				Ad26 Vector expressing the glycoprotein of the ebola virus mayinga variant [Ad26.ZEBOV]				Crucell Holland		
				2.Modified vaccinia ankara – bavarian nordic				B.V,		
				vector expressing the glycoproteins of ebola virus, sudan virus and marburg virus and the				Represented by Janssen	Approved but sponsor withdrew	
147 ZE	EBOV			nucleoprotein of tai forest virus [MVA-BN- Filo]/Vaccine	7th January 2015	Professor Fred Binka	OCRC, Hohoe	Pharmaceutica (Pty) Ltd	conduct N/A	
147				the ebola virus mayinga variant [Ad26.ZEBOV	7 ti bandary 2015	1 Tolessor Fred Birka		Crucell Holland	INA	
				Modified vaccinia ankara – bavarian nordic vector expressing the glycoproteins of ebola				B.V, Represented by		
				virus, sudan virus and marburg virus and the				Janssen		
148 ZE	EBOV 2	II	Ebola	nucleoprotein of tai forest virus [MVA-BN- Filo]/Vaccine	6th April 2015	Professor Fred Binka	OCRC, Hohoe	Pharmaceutica (Pty) Ltd	Application withdrawn N/A	
					·			. ,,		
							Noguchi Memorial Institute For Medical	General	Application Withdrawn N/A	
149 HY	YDRANON	l.		Hydranon solution	1st March 2008	Prof. David Ofori-Adjei	Research	Technology 1llc	N/A	
						1. Dr. Isaac Osei	Navrongo Health Research Centre	Janssen-Cilag		
								International NV		
						2. Dr. Samuel Abora	Upper East Regional Hospital	(Sponsor) represented by		
						3. Dr. Fred Adomako –	Kumasi Centre for Collaborative	Clinical	Application Withdrawn	
150 SA	ALIF,	IIIb	HIV	1.TDF/FTC/RPV 2.TDF/FTC/EFV/Vaccine	4th September 2013	Boateng	Research	Research Africa Ltd.	N/A	
								University of		
NO	OGUCHI SCD						Medical Research 2. College	Pittsburg, Representative:	Application Withdrawn	
151		IL.	Cialda Call Diaga	NVX-508/ Allopathic	1st May 2017	Amma Twumwaa Owusu Ansah	of Health Sciences 3.University of Ghana	Amma Owusu- Ansah, MD	N/A	
151		ID	Sickle Cell Disea	NVA-506/ Aliopatric	TSUMAY 2017	Arisari	Gnana	Alisali, MD		
										To address the gap in proteinuria measurement solutions, LifeAssay Diagnostics
								Emily Stephanie Zobrist, PATH,		(LAD) has developed and commercialized a low-cost PrCr urine dipstick that has shown goodlaboratoryand clinical performance and high usability within antenatal care
							Ridge Hospital,	2201 Westlake		(ANC)settings in previous studies. There is a need for further evidenceon the clinical
152 DE	RCR SPOT	Phase II	Procelomocio	PRCR Spot/Medical device	15th March 2021	Dr. Hannah Brown Amoakoh	Korlebu Teaching Hospital, Koforidua Regional Hospital	Avenue, Seattle, WA 98121, USA		utility and operational fit of the LAD Test-it™ PrCr test to inform policy recommendation for its use in Ghana and other LMIC settings.
132 PF	NON OF UT	i ilase II	гесыаттрыа	Tron oponivieuloai device	TOUT WATCH ZUZ I	AIIOAKUII	Trogional Fluspital	*** 30 12 1, USA	Оролзон	recommendation for its use in Oriana and other EMIC settings.
SA	AR97276A_SANOFI							Sanofi Aventis	Application Withdrawn by	
153		II	Malaria	SAR97276A/Allopathic	1st October, 2008	Prof. Seth Owusu-Agyei	Navrongo Health Research Centre	Recherche & Developpement	Sponsor before approval	
										General objective is to evaluate effects of polyphenol-rich cocoa as adjuvant therapy in
										COVID 19 patients. Specific objectives:  1. to determine the effects of natural polyphenol-rich natural cocoa powder (5 % v/w)
										(as adjuvant therapy) on symptom resolution and illness duration in COVID-19 patients
				Polyphenol-rich natural cocoa powder						to determine the effects of natural polyphenol-rich natural cocoa powder (5 % v/w)     on selected markers of coagulopathy in COVID-19 patients
PC	OLYPHENOL-RICH			Food supplements						to determine the effects of natural polyphenol-rich natural cocoa powder (5 % v/w) on virologic clearance COVID-19 patients
CC	OCOA POWDER						Ga East Municipal Hospital, Ghana	Ghana Cocoa	to unresponsiveness of	4. to determine the effects of natural polyphenol-rich natural cocoa powder (5% v/w)
154 TF	RIAL	Phase III	Covid-19	Oral	10th January 2022	Prof. George Obeng Adjei	Infectious Disease Centre	Board	applicant,, 4 Months	on disease prognosis COVID-19 patients

155	вемри	Phase II	Hyppthermia in Infants	Bempu Bracelet Medical device	2nd November, 2020	Mr. Prince Owusu	*Achimota General Hospital     *Greater Accra Regional Hospital     *Eastern Regional Hospital     *Korde-Bu Teaching Hospital     *Central Regional Hospital     Princess Marie Luis Children Hospital	Center for learning and childhood development	Application closed by FDA due to unresponsiveness of applicant,	To determine the accuracy of the bracelet in identifying hypothermia and evaluate its effect on Kangaroo Mother Care (KMC) practices and neonatal health outcomes in Ghana.  To assess the acceptability of the bracelet in Health providers and caregivers of Low Birth Weight (LBW) Infants by conducting qualitative in-depth interviews.  Determine the accuracy of the BEMPU bracelet in classifying hypothermia in the clinical setting.  Evaluate the impact of the bracelet
156	INO-9112 COVID 19	Phase I	Covid-19	INO-4800 followed by Electroporation (EP)     NO-4800 + INO-9112 followed by     Electroporation (EP)/ Vaccine	30th June 2022	Dr. Kwadwo Ansah Koram	Noguchi Memorial Institute for Medical Research, University of Ghana, Legon	Inovio Pharmaceuticals	Application closed by FDA due to unresponsiveness of applicant, 15 Months	The overall purpose of this clinical trial is to identify a booster dose of INO-4800 or INO 4800 plus INO-9112 given 6 to 12 months following primary vaccination with an approved or authorized mRNA vaccine for future development.
157	POST MASTECTOMY PAIN RELIEF			Erector Spinae block using bupivacaine/ Local anasthetics	2nd December 2021	Dr. Nana Addo Boateng	Komfo Anokye Teaching Hospital (KATH)	Self-Funding	Application closed by FDA due to unresponsiveness of applicant	General objective: The main objective of the study is to determine the postoperative analgesic effect of Erector Spines Plane (ESP) Block after mastectomy.  Specific objectives: 1. To compare the total morphine consumption within 24 postoperative hours between patients receiving ESP block with bupivacaine and ESP block with saline for mastectomy at the Komfo Anokye Teaching Hospital, Kumasi, Ghana. 2. To compare the numeric rating score at 2,4,6,12 and 24 hours between patients receiving ESP block with bupivacaine and ESP block with saline for mastectomy at the Komfo Anokye Teaching Hospital, Kumasi, Ghana. 3. To compare the time to the first request of rescue analgesia between patients receiving ESP block with bupivacaine and ESP block with saline for mastectomy at the Komfo Anokye Teaching Hospital, Kumasi, Ghana. 4. To compare patients satisfaction within the 24-hour postoperative analgesia between patients receiving ESP block with saline for mastectomy at the Komfo Anokye Teaching Hospital, Kumasi, Ghana.
158	SMAART-II	Phase III	STROKE	A polycap capsule contains Ramipril 5mg, Atenolol 50mg, Hydrochlorothiazide 12.5mg, Simvastatin 20mg, Aspirin 100mg.	16th August 2023	Dr. Fred Stephen Sarfo	Komfo Anokye Teaching Hospital (KATH)	University of California, San Francisco	Application closed by FDA	To deploy a hybrid study design to:  • firstly, demonstrate the efficacy of a polypill (Polycap ®) containing fixed doses of anthypertensives, a statin, and antiplatelet therapy taken as two capsules, once daily orally in reducing composite vescular risk over 24 months vs. usual care among 680 recent stroke patients encountered at 12 hospitals in Ghana.  - Secondly, SMAART II seeks to develop an implementationstrategy for routine integration and policy adoption of Polypill for post-stroke cardiovascular risk reduction in an under-resourced system burdened by suboptimal care and outcomes.
159	LETICIA	Phase II	Aneamia	LETICIA protocol diet (provided by study)     2.3-Fer syrup 3. Usual or Typical diet/ Food supplement	30th August, 2019	Dr. Lawrence Osei-Tutu	Agogo Presbyterian Hospital	Dr. Lawrence Osel-Tutu	Application closed by FDA since Sponsor/Pl failed to start study after approval	Iron deficiency is the most common nutritional deficiency worldwide and an important public health problem in Low and Middle Income Countries (LMICs). Causes of anemia in LMICs like Ghana are usually multifactorial including malaria, hemolytic anemias, and chronic blood loss from chronic parasitic infections including schildscomissis and hookworm. Factors accounting for inadequate supplies of dietary iron and micronutrients include poverty, a lack of nutritional supplementation, and food taboos. Anemia may result when iron deficiency is severe, after the body's iron stores are depleted and supply to the bone marrow is limited. This proof of concept study is to determine whether hospitalized children 6-59 months old who presented with moderate-to-severe anemia and given a combination of iron-rich food and standard iron replacement therapy (the intervention group) will demonstrate a greater final hemoglobin (Ho) concentration after two weeks compared to participants of similar characteristics in the control group who will receive oral iron supplementation in addition to their usual diet.
160	TENOFOVEK BE	Bioequivalenc		Trenofovek (tenofovir) 300mg film coated tablets 2. Viread (tenofovir) 300mg/Allopathic	11th September 2015	Prof. Seth Owusu Agyei     Dr. Kwaku Poku Asante	Kintampo Health Research Centre	Danadams Pharmaceuticals Industry Limited, Accra-Ghana	Application closed by FDA since Sponsor failed to start study 3 years after approval.	
161	ELDON CARD NYN	Feasibility stu		Eldon card 2. Standard laboratory method/Medical device	10th November 2015	Prof. Samuel Ameny Obed	Korle Bu Teaching Hospital, Accra.	Center for Global Child Health, Hospital for sick Children.	Incomplete CTA; Application closed by FDA.	

16	12 AX-100 HIVI		HIV	1.AX-100lmmun 2.AX-100lmmunPlus	9th december 2014	Dr. Kwaku Poku Asante	Kintampo Health Research Centre	Neopharmacie Limited , Germany	Incomplete CTA; Application closed by FDA.	
16	4P	III	Pregnancy Induced Hypertension and Preeclampsia	Polypil/Allopathic	9th August 2013	Dr. Emmanuel Kwabla Srofenyoh     Dr. Patrick Frimpong	Ridge Hospital Accra La General Hospital	Julius Centre for Health Sciences and Primary Care, University Medical Centre Utrecht, The Netherlands	Incomplete CTA; Application closed by FDA.	
16	INVACT	III	Malaria		13th may 2016	Prof. Kwadwo Ansah Koram	Noguchi Memorial Institute For Medical Research	Global Emerging Infections Surveillance and Response System of the US Armed Forces Health Surveillance Center		
									Incomplete CTA; Application closed by FDA.	
16	55 INSUGENIV	Phase IV	Diabetes	Insugen/Hormone	17th december 2013	N/A	Korle-Bu Teaching Hospital	BIOCON LTD	1071	
16	i6 AIM-LVRNA009	Phase IVIII	Covid-19	SARS-CoV-2 mRNA vaccine (LVR     Saline Placebol/vaccine	21st June 2022	Dr. Patrick Odum Ansah	1. Navrongo Health Research Centre 2. Kumasi Centre for Collaborative Research Health Research 3.Dodowa Health Research Centre 4. Kintampo Health Research Centre 5. Ghana Infectious Disease Centre 6. Korle Bu Teaching Hospital (KBTH)	AIM Vaccine Co. Ltd,	Not Approved,17-24 months.	Primary efficacy objective: To evaluate the protective efficacy of LVRNA009 (50 µg) in the prevention of first episodes of virologically-confirmed symptomatic cases of COVID-19 of any severity occurring from 14 days after 2nd dose in the initial set of vaccination in SARS-CoV-2 naive participants
	MYCOPIROX_LAGR AY		mixed Infection Vaginitis in					Lagray Chemical	Not Approved N/A	
16	57	Phase IV	Females	Mycopirox Vaginal cream	15th june 2010	Dr. Luitgard Darko		Company, Ltd.		
				Tobemstomig, Nab-Paclitaxel, Pembrotzumab		Dr. Hannah Naa Gogwe		F. Hoffmann-La	Study terminated by sponsor, 18	Primary Objective:
16	TNBC STUDY	Phase IIa	Breast Cancer	Monoclonal Antibody	28th December 2023	Ayettey Anie	Korle-Bu Teaching Hospital	Roche Ltd	months	□ To evaluate the immunogenicity to tobemstomig

169	VR-AD-1005 STUDY	Phase II	Cholera	VR-AD-1005/Aliopathic drug	1st July 2021	Dr. Ernest Kenu	Pentecost Hospital, Madina, Madina Polyclinic –	Vanessa Research Holdings, Inc.,	Study terminated by the sponsor due to safety issues 1 year 2 months	To assess the efficacy and safety of VR-AD-1005 for the treatment of acute diarrhea in cholera in combination with standard rehydration treatment with or without antibiotics (as indicated by WHO or other applicable guidelines) versus standard treatment alone. Efficacy is measured as reduction in stool output and/or duration of diarrhea between the start of treatment until final diarrheal stool before recovery or end of study treatment (treatment duration 120 hours).
170	ANTIPSYCHOTIC STUDY	Phase IV	Antipsychotic Induced Movement Disoders	Omega 3 Fish Oil Food supplement Oral	15th December 2021	Debrah Akosua Bema	Accra Psychiatric Hospital	Dr. Sammy Ohene P. O. Bu	Study terminated by sponsor due to safety issues, 29 Weeks	The primary objective of this study is to determine the use of once daily dose of 1000mg omega 3 fish oil as a clinically effective and safe intervention for reducing the burden associated with antipsychotic induced movement disorders. Secondary:  To determine the demographic and clinical characteristics of psychiatric patients with antipsychotic induced movement disorder.  To determine the efficacy of omega 3 supplementation in relieving the symptoms of AIM disorders To evaluate the impact of omega 3 supplementation on the clinical outcomes of psychosis, cognitive function and quality of life! adherence of participants. To determine the correlations between the demographic and clinical parameters and the outcomes of therapy To understand the experiences of patients who have used other complementary and alternative medicines aside omega 3 fish oil as adjunct to conventional therapy, in an attempt to be free from their symptoms
17	STAND	Phase III	Sickle Cell Disease	1.CRIZANLIZUMAB 2.PLACEBO/ Monoclonal antibody	30th September, 2019	1.Dr. Yvonne Dei Adomakoh 2.Dr. Vivian Paintsil	Ghana Institute of Clinical Genetics, Korle-Bu. Sickle Cell Office Directorate of Child Health.	Novartis Pharma AG	Study terminated by FDA due to safety issues. Yet to submit the final report. 8 years 5 months	Sickle cell disease (SCD) is a genetic blood disorder, caused by a single missense mutation in the $\beta$ -globin gene, progresses into a systemic disease. Vaso-occlusion is the hallmark of SCD and can lead to serious acute and chronic complications. Extensive preclinat at at has established P-selectin as a key mediator of VOC in SCD and suggest that its blockade or genetic absence of P-selectin decreases or eliminates its interactions with its ligands, thereby reducing vaso-occlusion. Crizanlizumab is a monoclonal antibody that binds to P-selectin preventing it interactions with its ligands. The purpose of this study is to compare the efficacy and safety of 2 doses of crizanlizumab (5.0 mg/kg and 7.5 mg/kg) versus placebo in adolescent and adult SCD patients (12 years and older) with history of VOC leading to healthcare visit.
172	ANTICOV	Phase III	Covid-19	Nitazoxanide, Ciclesonide, Paracetamol, Ivermectin, Artesunate Amodiaquine (ASAQ) Allopathic drug Oral	15th July, 2020	John Humphrey, AMUASI	Komfo Anokye Teaching Hospital	•Bernhard Nocht Institute for Tropical Medicine	Study terminated by sponsor due to safety issues and yet to submit Final report ,24 Months	The purpose of this study is to compare the efficacy of alternative treatment strategies versus control on the risk of progression to severe respiratory disease. As there is no validated animal model for COVID-19, the efficacy of any potential treatment remains speculative beyond what is known about their pharmacokinetic and in-vitro data. Several repurposed drugs are currently being tested in severe cases or as prophylaxis, and the results may become available by the time the present study is initiated. At the same time, a number of other drug candidates are being evaluated for in-vitro efficacy or in smal proof-of concept studies. 13 in view of the rapidly evolving landscape in Africa, it was decided to select an adaptive design for the study in order to allow for the flexibility of adding or dropping arms or adjusting the randomisation ratio based on the data as it becomes available. Additionally, given that the control arm in the study may not be acceptable in some countries, it was decided to adopt a master platform-based approach to be allow for integration of data from all sites in the interim analyses, irrespective of their ability to have randomised patients in all treatment arms.
	COVID 19 CHO- CELL(TERMINATED)			1.Recombinant two-component COVID-19 vaccine (CHO cell) 2. ReCOV Placebol/Vaccine	16th November 2021	Dr. Patrick Ansah	Dodowa Health Research Centre     Navorongo Health Research Centre	Jiangsu Recbio Technology Co., Ltd.		1.To evaluate the safety and reactogenicity of the recombinant two-component COVID-19 vaccine (CHO cell) (ReCOV for short) in adults aged 18 years and older.  2. To evaluate SARS-CoV-2 neutralizing antibody of ReCOV on Day 14 after 2 doses vaccination in adults aged 18 years and older.  3. To evaluate the efficacy of ReCOV in preventing RT-PCR confirmed symptomatic COVID-19 in adults aged 18 years and older.  4. To evaluate the safety and reactogenicity of ReCOV in adults aged 18 years and older.
174	MoRiOn	Phas II	Onchocerciasis	1.Rifanpentine (Priftin®) 2.Moxifloxacin (Avelox®) 3.Doxycycline/Vaccine	28th April, 2017	Prof. Alexander Yaw Debrah	1.Enchi Government Hospital     2.Communities of Aowin/Suaman     District W/R	Kumasi Centre for Collaborative Research in Tropical Medicine	Study terminated by sponsor due to safety issues. Yet to submit Final report 15 months	Onchocerciasis is caused by the parasite Onchocerca volvulus. More than 37 million people are estimated to be infected with 0. Volvulus worldwide. The current therapeutic strategy relies on annual mass drug administration (MDA) based on the drug donation program for Ivermectin. Ivermectin is mainly microfilaridatial and after a few months female worms resume MF production levels high enough for transmission. Therefore, safe microfilaridatid drugs are needed to reach the goal of elimination. The study aims to show efficacy (Wolbachia depletion) of combination Rifapentine plus Moxfilocaria using immunohistology compared to no treatment and treatment with Doxycycline.

1	COVID-19 5 MOUTHWASH	Phase III	Covid-19	Corsodyl Mouthwash 2.Wokadine mouthwash 3.Hydrogen Peroxide mouthwas	6th September 2021	Dr. George Boateng Kyei	Noguchi Memorial Institute for Medical Research	Dr. George Boateng Kyei	Study terminated by sponsor due to safety issues. Yet to submit Final report 1 year 6 months	To investigate how long it takes for SARS-CoV-2 asymptomatic or presymptomatic persons to shed viable virus. It also seeks to evaluate among these patients the effect of a one-time mouth rinse on the detectable viral load of SARS-CoV-2 and to determine how long it takes for SARS-CoV-2 with load to remain low after using the mouth rinse.
1	IMR-SCD	Phase lib	Sickle Cell Disease	1.IMR-687 2.IMR-687 Placebo/Allopathic	13th August 2020	1. Dr. Seyram Kaali 2. Dr. Olayemi Edeghongon	•Korle-Bu Teaching Hospital •Kintampo Health Research Centre	IMARA Inc.	Early termination by Sponsor due to safety issues 1 Year 7 Months	This is a phase 2b, randomized, double-blind, placebo-controlled, multicenter study of subjects aged 18 to 65 years with SCD (HbSS, HbSB0 thalassemia, or HbSB+ thalassemia) to evaluate the safety and efficacy of the PDE9 inhibitor, IMR-687 does of 23.0 to 34.5 mg/kg and 34.5 to 56.7 mg/kg, in a relevant model of anemia (Hbbth1/th1 mince), oral administration of IMR-687 or 30 days at 30 mg/kg/day (human equivalent dose of 2.4 mg/kg/day) or 60 mg/kg/day (human equivalent dose of 4.9 mg/kg/day) increased RBCs and Hb, and reduced refuculocytes. The degree of these changes was dose dependent, with statistically significant improvement at the higher dose of 60 mg/kg, in addition, IMR-687 at 60 mg/kg improved erythrobalst differentiation, suggesting a role for this compound in the improvement of ineffective erythropoiesis, a problem in a number of hemoglobin disorders
1	HESTIA4	Phase I	Sickle Cell Disease	Ticagrelor/ Allopathic	16th May, 2018	Dr. Patrick Ansah     Dr. Catherine Segbefia     Dr. Kokou Hefoume Amegan-Aho	Navrongo Health Research Centre     Korte-Bu Teaching Hospital     Volta Regional Hospital	AstraZeneca AB	Study termination due to safety issues 31 Months	Complications of sickle cell disease (SCD) occur very early in life. Painful crises first appear in the fingers and toes (dactylitis) in very young children prior to their first birthday. In addition to painful crises occurring in the very young, SCD can affect organ function early in life. Loss of spenic function begins as early as 5 months of age with associated increase in infection risk. Stroke risk begins at age 2. Given the early onset of symptoms and complications of this discorder, therapies for SCD should be targeted at children, including the very young. There is a need to first establish the pharmacokinetics (PK) of ticagretor in this age group to allow for modelling or extrapolation in this population.  This goal of the study is to evaluate PK data in the 0-2 year old population in order to way for further studies and ultimately use of ticagretor in this youngest population.
			Dicodo	Todgiston / mopume	Tour may, 2010			71000000000712		
1	TADO	III	Sickle Cell Disease in Pediatrics	Prasugrel/Allopathic	20th may 2013	Prof. Tsiri Agbenyega Dr. Catherine Idara Segbefia	Malaria Research Center, Agogo Korle-Bu Teaching Hospital, Accra – Korle Bu	Eli Lilly and Company Indianapolis	Prematurely terminated 24 months due to safety issues	
1	WOMAN '9	Ш	Postpartum Hemorrhage	Tranexamic acid(cyklokapronr injection)/ Allopathic	10th sept 2009	Dr. Opare Addo Henry Sakyi     Dr. Kwadwo Asamoah Nyarko-Jectey     Dr. Chris Opoku Fofie     Dr. Chris Bawa	Ashanti Mampong Municipal Hospital     Komfo Anokye Teaching Hospital	Clinical Trials Unit, London School of Hygiene and Tropical Medicine	Terminated by Sponsor Prematurely ended due to safety issues	
1	NEOVITA	III		Vitamin A		Dr. Sam Newton	Kintampo Health Research Centre	PATH	Premature Termination 36 Months due to safety issues	
	PROBIOTIC (MILD			Probiotic (Lactobacillus reuteri) Food supplement					FDA DISSOCIATED itself from any data or findings from the study due to violation of its	Aim To determine the therapsultic effects of probiotics in mild cognitively impaired individuals (MCI) at Korle-Bu Teaching Hospital. Specific objectives * To determine the bicavaliability of probiotics in mild cognitive individuals at Korle-Bu Teaching Hospital. * To determine the clinical effects of probiotics in mild cognitively impaired individuals at Korle-Bu Teaching Hospital. * To determine the molecular effects of probiotics in mild cognitively impaired individuals at Korle-Bu Teaching Hospital. * To determine the molecular effects of probiotics in healthy controls at Korle-Bu Teaching Hospital. * To determine the molecular effects of probiotics in healthy controls at Korle-Bu Teaching Hospital. * To determine the bicavaliability of probiotics in healthy controls at Korle-Bu Teaching
1	COGNITIVE IMPAIRMENT)	Phase I	Mild cognitive impairment	Oral	14th April 2023	Michael Quansah	Korle-Bu Teaching Hospital (KBTH)	University, Australia	guidelines for conducting clinical trials. 6 Months	Hospital.

								Study ended, FDA
								DISSOCIATED itself from any
								data or findings from the study due to violation of its guidelines
CALLASCOPE								for conducting clinical trials.
*			Pocket Colposcope (CALLASCOPE)/Medical			Ridge Hospital, Korle-Bu Teaching	Duke Global	3 months
182	ii	Cervical cancer	device 1	12th February 2019	Dr. Emmanuel Srofenyoh	Hospital	Health Institute	
								FDA DISSOCIATED itself from
						Hohoe Health Research Centre		any data or findings from the study due to violation of its
HOHOE			1.Dihydroartemisinin 2.Piperaquine oral				Malaria Capacity	guidelines for conducting clinical
ANTIMALARIAL			tablets 3.Artesunate 4.			Research Centre, Hohoe Municipal	Development	trials.
			Sulfamethoxypyrazine. 5.			Hospital, Ghana, Ghana Health	Consortium	7 months
183	III	Malaria	Pyrimethamine oral tablets/Allopathic		Dr. Margaret Kweku	Service	(MCDC 1. University of	Not Approved, FDA
							Ghana School	DISSOCIATES itself from any
			Azithromycin ,Injection Benzathine Penicillin				of Public Health	data or findings from the study
YAWS							2. World Health	due to violation of its guidelines
YAWS			Allopathic Drug		Dr. Cvnthia Kwakve-		Organization  3. Ghana Health	for conducting clinical trials.
184	III	Yaws	Oral		Maclean	Ga West District	Service, Ga	
								FDA DISSOCIATED itself from
			GMZ2 candidate malaria vaccine					any data or findings
405 CM7 0" / "		Malaria	Venning	10th average 2010	Dr. Fronk About	Navrongo Health Research Centre,	Statens Serum	27 onlhs
185 GMZ 2II / III		iviaiana		19th august 2010	Dr. Frank Atuguba	ivaviongo.	institute	
			Barley beta glucan					FDA DISSOCIATED itself from
			Food supplement				Best	any data Findings
		Cholesterol						N/Á
186 CEREBETA		concentration	Oral 1	13th may 2016	Mrs. Rose 1. Odotei Adjei	Suntreso Government hospital	Technologies	
							WORLD	
AQUAMAT			Artesunate, Quinine				HEALTH	EDA DIGOGGIATED N. M
107		Molorio	Allopathic 1	10th october 2012	Prof. Tsiri Agbenyega	Komfo Anokye Teaching Hospital	N	FDA DISSOCIATED itself from any data Findings
101	<b></b>	ivididila	/ mopulatio	10111 0010 001 2012	1 Tot. Tom / tgborryoga	Troming Funday Free Pilan		an) data i mungo
								FDA DISSOCIATED itself from
								any data or findings from the
						Ayensuanor District		study due to violation of its
AZI4YAWS						West Akyem Municipality		guidelines for conducting clinical
			A - di conservir			2. I to oct any off marriograms	World Health	
AZI4YAWS			Azythromycin			3. Upper West Akyem	Organization,	trials.
188	III	Yaws	Azythromycin Allopathic 2	23rd April 2015	Prof. Adu Sarkodie	3. Upper West Akyem		
188	III	Yaws	Azythromycin Allopathic 2	23rd April 2015	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
188	III	Yaws	Azythromycin Allopathic 2	23rd April 2015	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
188	III	Yaws	Azythromycin Allopathic 2	23rd April 2015	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
AZI4 TAWS	111	Yaws	Azythromycin Allopathic 2	23rd April 2015	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
AZIATAWS	111	Yaws	Allopathic 2	zoru zpri zoro	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
188	111	Yaws	Allopathic 2	23rd April 2015 SHORT AND DETAILED NAMES OF	Prof. Adu Sarkodie	3. Upper West Akyem	Organization,	trials.
188	A strategy to a	Yaws	Altopathic	SHORT AND DETAILED NAMES OF		Upper West Akyem     Nkwanta North District	Organization, Geneva - Switzerland	trals. 12 months
188	A strategy to r	Yaws	Allopathic 2	SHORT AND DETAILED NAMES OF		Upper West Akyem     Nkwanta North District	Organization, Geneva - Switzerland	trals. 12 months
188			Allopathic 2	SHORT AND DETAILED NAMES OF Maternal Mortality by 50% or m	nore Polypill for the Prever	Upper West Akyen     Nkwanta North District      Nkwanta North District  ntion of Pregnancy Induced Hypertension	Organization, Geneva - Switzerland	trals. 12 months
1 4P ABDOV COVID 19 2 TRIAL	A randomized	, double-blind, po	Allopathic 2  ons of Hypertensive disorders in Pregnancy and M stillve-controlled Phase III clinical trial to evaluate th	short and Detailed names or Maternal Mortality by 50% or m he efficacy and safety of SCT	nore Polypill for the Prever	Newanta North District     Newanta North District  Intion of Pregnancy Induced Hypertension  Beta/Delta/Omicron Variants S Trimer Variants  Strimer Variants S Trimer Variants S Trimer Variants  Seta/Delta/Omicron Variants S Trimer Variants  Seta/Delta/Delta/Omicron Variants S Trimer Variants  Seta/Delta	Organization, Geneva - Switzerland  and Preeclamps cocine) in population	tials. 12 months ia (4P) Trial
1 4P ABDOV COVID 19	A randomized	, double-blind, po	Allopathic 2	short and Detailed names or Maternal Mortality by 50% or m he efficacy and safety of SCT	nore Polypill for the Prever	Newanta North District     Newanta North District  Intion of Pregnancy Induced Hypertension  Beta/Delta/Omicron Variants S Trimer Variants  Strimer Variants S Trimer Variants S Trimer Variants  Seta/Delta/Omicron Variants S Trimer Variants  Seta/Delta/Delta/Omicron Variants S Trimer Variants  Seta/Delta	Organization, Geneva - Switzerland  and Preeclamps cocine) in population	tials. 12 months ia (4P) Trial
1 4P ABDOV COVID 19 2 TRIAL 3 ACTIVE TRIALS	A randomized A Phase 3, m	, double-blind, po	Allonathic 2  ons of Hypertensive disorders in Pregnancy and M ssitive-controlled Phase III clinical trial to evaluate the	SHORT AND DETAILED NAMES OF Atternal Mortality by 50% or m he efficacy and safety of SCT and antiviral effect of S-21762;	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in	Newanta North District     Newanta North District  antion of Pregnancy Induced Hypertension  Beta/Deta/Omicron Variants S Trimer Var non-hospitalized participants with COVIC	Organization, Geneva - Switzerland  and Preeclamps ccine) in population	tials.  12 months  ia (4P) Trial  in previously unvaccinated with COVID-19 vaccine and aged ≥18 years
1 4P ABDOV COVID 19 2 TRIAL	A randomized A Phase 3, m	, double-blind, po	Allonathic 2  ons of Hypertensive disorders in Pregnancy and M ssitive-controlled Phase III clinical trial to evaluate the	SHORT AND DETAILED NAMES OF Atternal Mortality by 50% or m he efficacy and safety of SCT and antiviral effect of S-21762;	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in	Newanta North District     Newanta North District  antion of Pregnancy Induced Hypertension  Beta/Deta/Omicron Variants S Trimer Var non-hospitalized participants with COVIC	Organization, Geneva - Switzerland  and Preeclamps ccine) in population	tials. 12 months ia (4P) Trial
1 4P ABDOV COVID 19 2 TRIAL 3 ACTIVE TRIALS 4 AIM-LVRNA009	A randomized A Phase 3, m A Global Multi	, double-blind, poulticenter, random	ons of Hypertensive disorders in Pregnancy and M stitive-controlled Phase III clinical trial to evaluate the sized, double-blind, 24-week study of the clinical arized, Blinded, Placebo-controlled Phase 2/3 Clinical arized ar	SHORT AND DETAILED NAMES OF Asternal Mortality by 50% or in the efficacy and safety of SCT and antiviral effect of S-217622 as Study to Evaluate the Effici	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in acy, Safety and Immunogen	A. Nicwanta North District     Nicwanta North District     The Control of Pregnancy Induced Hypertension Seta/Delta/Omicron Variants S Trimer Variant-hospitalized participants with COVID inicity of SARS-CoV-2 mRNA Vaccine (LV)	Organization, Geneva - Switzerland  and Preeclamps coine) in populatic 0-19  RNA009) for the	tials.  12 months  ia (4P) Trial  in previously unvaccinated with COVID-19 vaccine and aged ≥18 years
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1 4P ABDOV COVID 19 2 TRIAL 3 ACTIVE TRIALS 4 AIM-LVRNA009 5 AIMS	A randomized A Phase 3, m A Global Multi	, double-blind, poulticenter, random	ons of Hypertensive disorders in Pregnancy and M stitive-controlled Phase III clinical trial to evaluate the sized, double-blind, 24-week study of the clinical arized, Blinded, Placebo-controlled Phase 2/3 Clinical arized ar	SHORT AND DETAILED NAMES OF Asternal Mortality by 50% or in the efficacy and safety of SCT and antiviral effect of S-217622 as Study to Evaluate the Effici	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in acy, Safety and Immunogen	A. Nicwanta North District     Nicwanta North District     The Control of Pregnancy Induced Hypertension Seta/Delta/Omicron Variants S Trimer Variant-hospitalized participants with COVID inicity of SARS-CoV-2 mRNA Vaccine (LV)	Organization, Geneva - Switzerland  and Preeclamps coine) in populatic 0-19  RNA009) for the	tials.  12 months  ia (4P) Trial  in previously unvaccinated with COVID-19 vaccine and aged ≥18 years
1 4P ABDOV COVID 19 2 TRIAL 3 ACTIVE TRIALS 4 AIM-LVRNA009	A randomized A Phase 3, m A Global Multi African Investi	, double-blind, pout ulticenter, random -center, Random gation Of Mirasol	ons of Hypertensive disorders in Pregnancy and M stitive-controlled Phase III clinical trial to evaluate the sized, double-blind, 24-week study of the clinical arized, Blinded, Placebo-controlled Phase 2/3 Clinical arized ar	SHORT AND DETAILED NAMES OF Asternal Mortality by 50% or in the efficacy and safety of SCT and antiviral effect of S-21762; asl Study to Evaluate the Effici Efficacy Of Mirasol Treated F	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in acy, Safety and Immunogen resh Whole Blood For The I	13. Upper West Akyem 4. Nkwanta North District  antion of Pregnancy Induced Hypertension Seta/Delta/Omicron Variants S Trimer Var non-hospitalized participants with COVID icitly of SARS-CoV-2 mRNA Vaccine (LV Prevention Of Transfusion Transmitted M	Organization, Geneva - Switzerland  and Preeclamps coine) in populatic 0-19  RNA009) for the	tials.  12 months  ia (4P) Trial  in previously unvaccinated with COVID-19 vaccine and aged ≥18 years
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1 4P ABDOV COVID 19 2 TRIAL 3 ACTIVE TRIALS 4 AIM-LVRNA009 5 AIMS	A randomized A Phase 3, m A Global Multi African Investi Comparison c	, double-blind, po ulticenter, random -center, Random gation Of Mirasol	ons of Hypertensive disorders in Pregnancy and M ssilive-controlled Phase III clinical trial to evaluate th nized, double-blind, 24-week study of the clinical ar ized, Blinded, Placebo-controlled Phase 2/3 Clinical System For Whole Blood, Clinical And Biological	SHORT AND DETAILED NAMES OF Alaternal Mortality by 50% or made efficacy and safety of SCT and antiviral effect of S-21762: all Study to Evaluate the Efficacy Of Mirasol Treated Futheir efficacy against Onchoos	nore Polypill for the Prever V01E (A COVID-19 Alpha/E 2 compared with placebo in acy, Safety and Immunogen resh Whole Blood For The I	13. Upper West Akyem 4. Nkwanta North District  antion of Pregnancy Induced Hypertension Seta/Delta/Omicron Variants S Trimer Var non-hospitalized participants with COVID icitly of SARS-CoV-2 mRNA Vaccine (LV Prevention Of Transfusion Transmitted M	Organization, Geneva - Switzerland  and Preeclamps coine) in populatic 0-19  RNA009) for the	tials.  12 months  ia (4P) Trial  in previously unvaccinated with COVID-19 vaccine and aged ≥18 years
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12 AQUAMAT	
12 AQUAMAT	An Open Randomized Comparism of Artesunate versus Quinine in the Treatment of Severe Falciparum Malaria in African Children.
13 ARTIMIST	A Phase III, Randomized, Open Labelled, Active Controlled, Multicentre, Superiority Trial Of Artimistim Versus Intravenous Quinine In Children With Severe Or Complicated Falciparum Malaria, Or Uncomplicated Falciparum Malaria With Gastrointestinal Complications
14 ASAAP	A Multicentre Phase III Non-Inferiority Trial to Evaluate Safety, Tolerability and Efficacy of Artemether-Lumefantrine+Atovaquone-Proquanil Tri-Therapy Versus Artemether-Lumefantrine Bi-Therapy for the Treatment of Uncomplicated Malaria in African Children Aged 6 Months To 10 Years (ASAAP PROJECT)
15 ASTAWOL	The efficacy of Rifampicin 35mg/Kg/d plus Albendazole 400mg/d given for 7 or 14 days against Lymphatic Filariasis and Onchocerciasis- a randomized, controlled, parallel-group, open-label, phase II pilot trial
16 ATEA COVID 19	A Phase 3 Randomized, Double-Blind, Placebo-Controlled Study to Evaluate the Efficacy and Safety of Bennifosbuvir in High-Risk Outpatients with COVID-19
10 ATEA COVID 19	A Friase 3 Nationitized, Doduce-billo, Fisicator-Controlled Study to Evaluate the Emicary and Salety of Denninoscount in High-Yask Coupations with COVID-19
17 AVAREF	A Phase 3 double-blind, randomized, active comparator-controlled, group-sequential, multinational trial to assess the safety, immunogenicity and efficacy of a trivalent rotavirus P2-VP8 subunit vaccine in prevention of severe rotavirus gastroenteritis in healthy infants.
18 AX-100 HIV	A Double Blind Randomized Control Trial of AX-100 Immun (Liquid) and AX-100 Immun Plus Combination Among Adults Living with HIV In Ghana.
AZIDUS 19 ACECLOFENAC	An open label, balanced, randomized, two treatments, two periods, two sequences, single dose, crossover, relative bioavailability study of two different formulations of Aceclofenac tablets 100 mg (T1 & T2) of OA&J Pharmaceuticals Ltd, Ghana in healthy adult human subjects under fasting condition
AZIDUS 20 BUPRENORPHINE	An open label, balanced, randomized, two treatments, two periods, two sequences, single dose, crossover, bioequivalence study of Buprenorphine 16 mg Sublingual tablets of Wes Pharma Inc and Buprenorphine hydrochloride 8 mg (8 mg x 2 tablets) sublingual tablets of Hikma Pharmaceuticals USA Inc in healthy adult human subjects under fasting condition
AZIDUS 21 CEFUROXIME	An open label, balanced, randomized, two treatments, two periods, two sequences, single dose, crossover, bioequivalence study of Cefuroxime Axetil 500 mg Tablets of OA&J Pharmaceuticals Ltd, Ghana and Zinnat (Cefuroxime Axetil) 500 mg film-coated tablets of GlaxoSmithKline UK in healthy adult human subjects under fed condition
22 AZI4YAWS	Randomized Controlled Trial Comparing Efficacy of a Single Dose of Treatment of Yaws with 20mg/kg versus 30mg/kg of Azithromycin.
AZITHROMYCIN PLUS CHLOROQUINE 23 PHOSPHATE	Azithromycin Plus Chloroquine Phosphate versus Artemether-Lumefatrine for the Treatment of Uncomplicated Plasmodium falciparium Malaria in Children in Africa.
24 BEMPU	Hypothermia Prevention in low birth weight and preterm Infants
25 BILI-RULER	Improving community-based diagnosis of neonatal jaundice using a simple icterometer: The Bill-Ruler Study
26 BLMs4BU	SHORTENING BURULI ULCER TREATMENT: WHO RECOMMENDED VS. A NOVEL BETA-LACTAM-CONTAINING THERAPY – PHASE III EVALUATION INWEST AFRICA
27 BURULINOX	Evaluation of nitric oxide generating dressing (EDX) to improve management of buruli ulcer disease – a prospective randomized open-blinded end point.
28 BURULIRIFDACC	A randomized controlled trial to evaluate the effect of High Dose of Rifampicin and Dialkylcarbamoyl chloride (DACC)-coated dressings on outcomes in Mycobacterium ulcerans disease

29 CDA	A Multicenter, Randomized, Double Blind Study to Compare the Efficacy and Safety of CDA Versus Artemether-Lumefantrine in the Treatment of Acute Uncomplicated P. Falciparum Malaria in Children and Adults in Africa.
30 CDA2	A Multicenter, Randomized, Double Blind Study to Compare the Efficacy and Safety of CDA Versus Chlorproguanil-Dapsone in the Treatment of Acute Uncomplicated P. Falciparum Malaria in Children and Adults in Africa.
31 CEREBETA	Efficacy of Beta-Glucans from Barley and Maintenance of Normal Blood LDL-Cholesterol Concentrations: A Randomized Control Study in Ghana.
32 CPAP	Clinical Trial Evaluating the Difference in Mortality Rates in Children in Ghana Receiving Continuous Positive Airway Pressure (CPAP) Versus Those Who Do Not.
33 CRASH-2	A Large Randomized Placebo Controlled Trial, among trauma patients with or at risk of significant Haemorrhage, of the Effects of Anti-Fibrinolytic treatment on Death and Transfusion requirement
34 CALLASCOPE	Clinical Studies and in-Depth Interviews for Portable, low-cost and Speculum-Free Cervical Cancer Screening in Ghana
35 CECOLIN	Phase 3 Randomized, Active-Comparator Controlled, Open-Label Trial to Evaluate the Immunogenicity and Safety of Alternate Two-Dose Regimens of a Bivalent Human Papillomavirus (HPV) Vaccine (Cecolin®) Compared to a Licensed Quadrivalent HPV Vaccine (Gardasil®) in Healthy 9-14 Year-Old Girls in Low and Low-Middle Income Countries
CEPHEIDXPERT HIV-	
36 1	An Investigation to Evaluate the Performance of the Cepheid XpertR HIV-1 VL XC Test
37 CIELO	A Phase III, Randomized, Double-blind, Placebo-controlled, Multicenter Basket Study to Evaluate the Efficacy, Safety, Pharmacokinetics, and Pharmacodynamics of Satralizumab in Patients with Anti-N-Methyl-D-Aspartic Acid Receptor (NMDAR) or Anti-Leucine-Rich Glioma-Inactivated 1 (LGI1) Encephalltis
38 CLARITY AFRICA	CiLostAzol for pRevention of recurrent sTroke in Africa (CLARITY-AFRICA): A Phase III Randomized Clinical Trial
39 COPE TRIAL	Effectiveness and Acceptability of two models of an Insertable Vaginal Cup for Non-surgical management of obstetric fistula in Ghana: a hybrid type 1 randomized crossover trial
40 COVID ABDOV	A randomized, double-blind, positive-controlled Phase III clinical trial to evaluate the efficacy and safety of SCTV01E (A COVID-19 Alpha/Beta/Delta/Omicron Variants S Trimer Vaccine) in population previously unvaccinated with COVID-19 vaccine and aged ≥18 years* (COVID ABDOV).
CROWN 41 CORONATION	An international, Bayesian platform adaptive, randomized, placebo-controlled trial assessing the effectiveness of candidate interventions in preventing COVID-19 disease in healthcare workers
42 CHEETAH	Cluster Randomized Trial of Sterile Glove and Instrument Change at the Time of Wound Closure to Reduce Site Infection: A Trial In Low- And Middle-Income Countries (LMICs)
TE OTTEETTWT	Consider I frame of October Chine On the Bridge State Chine On Whole Checked the Industrial And I frame in Light State Chine C
43 COVID 19 CHO-CELL	A multicenter, randomized, double-blind, placebo-controlled Phase II/III trial to evaluate the efficacy, safety and immunogenicity of the recombinant two-component COVID-19 vaccine (CHO cell) in adults aged 18 years and older
COVID 19	
44 INTRANASAL SPRAY COVID 19	A Global, Multi-center, Randomized, Double-blind, Placebo-controlled Phase III Clinical Trial to Evaluate the Protective Efficacy and Safety of Influenza Virus Vector COVID-19 Vaccine for Intranasal Spray (DelNS1-2019-nCoV-RBD-OPT1) in Adults Aged 18 Years and Older
45 MOUTHWASH	Viral Shedding Dynamics and the Effect of Antimicrobial Mouthwashes on the Detection of SARS-CoV-2 in Ghana.
DIABETIC FOOT 46 CARE	Service Advanced Districts From Self-one Decreasing in Change & Frontier Management of Controlled Taiglight Services on the Management of Controlled Taiglight
47 DOLF_IDA	Family-oriented Diabetic Foot Self-care Programme in Ghana; A Feasibility Randomised Controlled Trial with nested qualitative interviews at the Komfo Anokye Teaching Hospital.  Safety and Efficacy of Combination Therapy with Ivermectin, Diethylcarbamazine and Albendazole (IDA) for Individuals with Onchooerciasis
48 DRAGON	Multicentre non-inferiority cluster randomised trial testing Disposable versus Reusable drApes and Gowns for green operating theatres
49 EBA	Double-Blinded, Placebo-Controlled Dosage-Escalation Study and Immunogenicity of EBA-175 RII-NG Malaria Vaccine Administered Intramuscularly in Semi Immune Adults
50 EBOLA Z	A Phase 2, Randomized, Observer-Blind, Placebo-Controlled, Multi-Country Study to Assess the Safety and Immunogenicity of a Single Inframuscular Dose of GSK Biologicals' Investigational Recombinant Chimpanzee Adenovirus Type 3 – Vectored Ebola Zaire Vaccine. (ChAd3-EBO-Z) (GSK3390107A), in Adults 18 years of age and older in Africa
EBOLA Z 51 (PAEDIATRIC)	A Phase 2, Randomizad, Observer-Blind, Placebo-Controlled, Multi-Country Study to Assess the Safety and Immunogenicity of a Single Intramuscular Dose of GSK Biologicals' Investigational Recombinant Chimpanzee Adenovirus Type 3 — Vectored Ebola Zaire Vaccine. (ChAd3-EBO-Z) (GSK3390107A), in children
, ,	1 to 17 years of age in Africa
52 EBSI-LSV	
	A Phase 1 Randomized, Blinded, Placebo Controlled, Dose-Escalation and Dosing Regimen Selection Study to Evaluate the Safety and Immunogenicity of rVSV-Vectored Lassa Virus Vaccine in Healthy Adults at Multiple Sites in West Africa
53 ELDON CARD	A Phase 1 Randomized, Blinded, Placebo Controlled, Dose-Escalation and Dosing Regimen Selection Study to Evaluate the Safety and Immunogenicity of rVSV-Vectored Lassa Virus Vaccine in Healthy Adults at Multiple Sites in West Africa  Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana
54 EMODEPSIDE	
54 EMODEPSIDE 55 ESM UBT	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana
54 EMODEPSIDE	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.
54 EMODEPSIDE 55 ESM UBT	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centire Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage
54 EMODEPSIDE 55 ESM UBT 56 FALCON	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 1/2/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBIT/XIAOMI	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 112/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBIT/XIAOMI 60 FLORAL STUDY FORTIFIED BUILLON 61 CUBES STUDY	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 112/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors  An open-label, multi-centre, rollover study to characterise long-term safety and efficacy of etavopivat in adults, adolescents and children who have sickle cell disease or thalassaemia and have completed a treatment period in an etavopivat study  Effect of household use of multiple micronutrient-fortified bouillon on micronutrient status among women and children in two districts in the Northern region of Ghana
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBIT/XIAOMI 60 FLORAL STUDY FORTIFIED BUILLON 61 CUBES STUDY 62 GARDASIL	Using Eklon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchoerea volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 1/2/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors  An open-label, multi-centre, roliover study to characterise long-term safety and efficacy of etavophyat in adults, adolescents and children who have sickle cell disease or thalassaemia and have completed a treatment period in an etavophyat study  Effect of household use of multiple micronutrient-fortified bouillon on micronutrient status among women and children in two districts in the Northern region of Ghana  Evaluation of Safety And Immunogenicity Of Gardasitm in Healthy Females Between 9 And 26 Years Of Age in Subsaharan Africa
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBITIXIAOMI 60 FLORAL STUDY FORTIFIED BUILLON 61 CUBES STUDY 62 GARDASIL 63 GBT021601-021	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 1/2/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors  An open-label, multi-centre, rollover study to characterise long-term safety and efficacy of tavopivat in adults, adolescents and children who have sickle cell disease or thalassaemia and have completed a treatment period in an etavopivat study  Effect of household use of multiple micronutrient-fortified bouillion on micronutrient status among women and children in two districts in the Northern region of Ghana  Evaluation of Safety And immunogenicity Of Gardasiltm in Healthy Females Between 9 And 28 Years Of Age in Subsaharan Africa  A Phase 2/3 Randomized, Multicenter Study of Osivelotor Administered Orally to Participants with Sickle Cell Disease and an Open-Label Pharmacokinetics Study in Pediatric-Participants with Sickle Cell Disease
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBIT/XIAOMI 60 FLORAL STUDY FORTIFIED BUILLON 61 CUBES STUDY 62 GARDASIL	Using Eklon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgical Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 1/2/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Filoviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors  An open-label, multi-centre, roliover study to characterise long-term safety and efficacy of etavophyat in adults, adolescents and children who have sickle cell disease or thalassaemia and have completed a treatment period in an etavophyat study  Effect of household use of multiple micronutrient-fortified bouillion on micronutrient status among women and children in two districts in the Northern region of Ghana  Evaluation of Safety And Immunogenicity Of Gardasiltm in Healthy Females Between 9 And 26 Years Of Age in Subsaharan Africa
54 EMODEPSIDE 55 ESM UBT 56 FALCON 57 FERROQUINE 58 FILOVIRUS 59 FITBITIXIAOMI 60 FLORAL STUDY FORTIFIED BUILLON 61 CUBES STUDY 62 GARDASIL 63 GBT021601-021	Using Eldon Card for Testing of Maternal and Newborn Blood Group in Comparison with the Standard Laboratory Method of Blood Group Testing in Accra, Ghana  A phase II, Randomised, double-blind, parallel – group trial to investigate Emodepside (BAY 44-4400) in subjects with onchocerca volvulus infection.  A Multi-Centre Prospective Trial on the Impact of the Introduction of Condom-Based Uterine Balloon Tamponade for Uncontrolled Postpartum Hemorrhage  Pragmatic Multicentre Factorial Randomized Controlled Trial Testing Measures to Reduce Surgicial Site Infection in Low and Middle Income Countries  Randomized Multicentre Study Evaluating the Safety and Activity of Ferroquine Associated with Artesunate versus a Positive Calibrator (Amodiaquine Associated with Artesunate) in African Adult Patients with Uncomplicated Malaria  A phase 1/2/3 study to evaluate the safety, tolerability, immunogenicity, and efficacy of vaccine candidates against (Floviruses) disease in healthy individuals at risk of (Filoviruses) disease.  Feasibility of a wireless monitoring system as an alternative to current bedside monitors  An open-label, multi-centre, rollover study to characterise long-term safety and efficacy of etavopivat in adults, adolescents and children who have sickle cell disease or thalassaemia and have completed a treatment period in an etavopivat study  Effect of household use of multiple micronutrient-fortified bouillon on micronutrient status among women and children in two districts in the Northern region of Ghana  Evaluation of Safety And immunogenicity Of Gardasiltm in Healthy Females Between 9 And 26 Years Of Age in Subsaharan Africa  A Phase 2/3 Randomized, Multicenter Study of Osivebtor Administered Orally to Participants with Sickle Cell Disease and an Open-Label Pharmacokinetics Study in Pediatric-Participants with Sickle Cell Disease

67 GBT440-038	An Open-Label Extension Study of Voxelotor Administered Orally to Participants with Sickle Cell Disease Who Have Participated in Voxelotor Clinical Trials
68 GMZ 2	Randomized, Controlled, Double-Blind, Multicentre Study To Evaluate The Efficacy, Safety And Immunogenicity Of GMZ2 Candidate Malaria Vaccine In Gabonese, Burkinabe, Ghanaian And Ugandan Children Aged 12-60 Months
69 HIBISCUS	A global phase 3, randomised, double-blind and placebo-controlled study evaluating the efficacy and safety of etavopivat in adolescents and adults with sickle cell disease
70 HESTIA4	A Multi-centre, Phase I, Open-label, Single-dose Study to Investigate Pharmacokinetics (PK) of Ticagrelor in Infants and Toddlers, Aged 0 to less than 24 Months, with Sickle Cell Disease
71 HESTIA3	A Randomised, Double-Blind, Parallel-Group, Multicentre, Phase III Study to Evaluate the Effect of Ticagrelor versus Placebo in Reducing the Rate of Vaso-Occlusive Crises in Paediatric Patients with Sickle Cell Disease
HOHOE 72 ANTIMALARIAL	A Phase III of the Assessment of the Efficacy, Tolerability and Ease of Administration of, Dihydroartemisinin Plus Piperaquine and and Artesunate Plus Sulfamethoxypyrazine Plus Pyrimethamine for preventing Malaria in Ghanaian Children
73 HOPE SCD	A Phase 3, Double-blind, Randomized, Placebo-controlled, Multicenter Study of GBT440 Administered Orally to Patients With Sickle Cell Disease
74 HOPE KIDS 2	A phase 3,Randomised,Double-Blind, Placebo-Controlled Study of Voxelotor(GBT440) in Pediatric Participants with Sickle Cell Disease.
75 HYDRANON	Hydranon® solution (GR-08) in healthy adult volunteers
76 IAVI C105	A Phase 2 Randomized, Double-Blinded, Placebo-Controlled Clinical Trial to Evaluate the Safety, Tolerability, and Immunogenicity of rVSV\(\Delta\)G-LASV-GPC Vaccine in Adults and Children Residing in West Africa
77 IMBRAVE 152	A phase III, randomized, double-blind, placebo-controlled, study evaluating Atezolizumab and Bevacizumab, with or without Tiragolumab, in patients with untreated locally advanced or Metastatic Hepatocellular Carcinoma
78 IMR-SCD-301	A Phase 2b Study to Evaluate the Safety and Efficacy of IMR-687 in Subjects with Sickle Cell Disease
79 INNOVATE	Phase 2/3 Randomized, Blinded, Placebo-Controlled Trial to Evaluate the Safety, Immunogenicity, and Efficacy of INO-4800, a Prophylactic Vaccine against COVID-19 Disease, Administered Intradermally Followed by Electroporation in Adults at High Risk of SARS-CoV-2 Exposure
80 INO-9112 COVID 19	Phase 1 Open Label, Randomized Study to Evaluate the Safety, Tolerability, and Immunogenicity of an Intradermal Booster Dose of INO-4800 alone or in combination with INO-9112 followed by Electroporation in Adults who Completed a Primary Immunization Series Against SARS-CoV-2 with mRNA Vaccines
81 INVACT	In Vivo Efficacy of Artemisinin Combination Therapy to Explore Laboratory and Parasitological Markers of Artemisinin Resistance in Uncomplicated Plasmodium falciparum Malaria in Ghana.
82 IPT & SP	Operational Research on Intermittent Preventive Treatment of Malaria in Infants (IPTi) with Sulfadoxine/Pyrimethamine (S/P)
83 INSUGEN	Post Market Surveillance Study of Insugen 30/70
	A Phase Ila observer-blind, randomized, controlled, age-de-escalation, single center interventional study to evaluate the safety, reactogenicity, and immune
84 INTS GMMA INOVIO – LASSA	response of the GVGH iNTS vaccine against S. Typhimurium and S. Entertitids, in adults, children and infants,
85 FEVER	Study to evaluate the safety, tolerability and immunogenicity of INO-4500 in Healthy volunteers
IRON 86 FORTIFICATION	Seasonal Impact Of Iron Fortification On Malaria Incidence In Ghanaian Children
87 IUMO	RANDOMISED CONTROLLED TRIAL: INTRAUTERINE MISOPROSTOL VERSUS SUBLINGUAL MISOPROSTOL IN THE PREVENTION OF POSTPARTUM HEMORRHAGE AT ELECTIVE CAESAREAN SECTION AT KORLE BU TEACHING HOSPITAL.
88 IVERMECTIN GH	Safety and Efficacy of Ivermectin in the Prevention and Management of COVID- 19 among Ghanaian Populations

89 KAE609	A Phase 2, Multi-Center, Randomized, Open - Label, Dose Escalation Study To Determine Safety Of single (QD) and Multiple (3QD) Doses Of KAE609, Given To Adults With Uncomplicated Plasmodium Falciparum Malaria
91 KALUMA	A randomized, open-label, multicenter study to compare efficacy, safety and tolerability of KLU156 with Coartem® in the treatment of uncomplicated Plasmodium falciparum malaria in adults and children ≥ 5 kg body weight followed by an Extension phase with repeated KLU156 treatment
92 KANGAROO CARE	Enhancing the Survival of Low Birthweight Infants in Low Resource Settings Using an Implementation Science Approach
93 KNC 19(NIBIMA)	Repurposing the aqueous Extract of Cryptolepis for Covid-19 therapy
94 LEDoxy	Doxycycline 200mg/d vs. 100mg/d for 6 weeks to improve filarial lymphedema - a multinational, double-blind, randomized, placebo-controlled trial.
95 LETICIA	Combination Food-Based And Supplemental Iron Replacement Therapy For Children With Moderate-To-Severe Anemia In A Rural Ghanaian Setting: A Proof-Of-Concept Study
96 LIVZON	A Global, Multi-Center, Randomized, Double-Blind, Placebo-Controlled, Phase III Clinical Study to Evaluate the Efficacy, Safety, and Immunogenicity of Recombinant SARS-CoV-2 Fusion Protein Vaccine (V01) in Adults Aged 18 Years and older.
97 MAL 047	Randomized, Controlled, Partially-Blind Study Of The Safety And Immunogenicity Of Glaxosmithkline Biologicals' Candidate Plasmodium Falciparum Vaccines RTS,S/AS02D And RTS,S/AS01E, When Administered IM According To A Three Dose Schedules In Children Aged 5 To 17 Months Living In Ghana.
98 MAL 050	Randomized, Open, Controlled Study Of The Safety Of The And Immunogenicity Of GSK Biologicals' Candidate Plasmodium Falciparium Malaria vaccine RTS, S/AS01E when incorporated into an expanded program on immunization (EPI) regimen that includes DTPWHEPB/HIB.OPV, Measles and yellow fever vaccination in infants living in malaria- Endemic Regions- 050
99 MAL 055	Double Blind (Observer Blind), Randomised, Controlled Multicentre Study To Evaluate In Infants And Children, The Efficacy Of RTS,S/AS10E Candidate Vaccine Against Malaria Disease Caused By P. Falciparium Infection Across Diverse Malaria Transmission Settings In Africa
100 MAL 063	Randomized, Open, Controlled Study To Evaluate The Immune Response To The Hepatitis B Antigen Of The RTS,S /AS01E Candidate Vaccine, When Administrated As Primary Vaccination Integrated Into An EPI Regimen To Infants Living In Sub-Saharan Africa
	Phase Illb randomized, open, controlled, multi-center study to evaluate the immunogenicity and safety of the RTS,S/AS01E candidate malaria vaccine, when administered as primary vaccination at 6, 7.5 and 9 months of age with or without co-administration of measles, rubella and yellow fever vaccines followed by an
101 MAL 073	RTS_S/AS01E booster vaccination 18 months post Dose 3, to children living in sub-Saharan Africa  Phase lib Randomized, Open-Label, Controlled, Multi-Centre Study of the Efficacy, Safety and Immunogenicity of GSK Biologicals' Candidate Malaria Vaccine RTS_S/AS01E Evaluating Schedules with or without Fractional Doses, early Doses, in Children 5-17 Months of age Living in Sub-Saharan
102 MAL 094	Phase in Kandomized, Open-Labet, Controlled, Multi-Centre Study of the Emicacy, Sately and immunogenicity of GSK biologicals Candidate Malaria Vaccine KTS, S/ASUTE Evaluating Schedules with of without Fractional Doses, early Dose 4 and yearly Doses, in Children 5-17 Months of age Living in Sub-Sanaran Africa.
103 MALHELMINTHS	Evaluating the effectiveness and cost-effectiveness of integrating mass drugadministration for helminth control with seasonal malaria chemoprevention in Ghanaian children
104 MDGH-MOX-1006	An open-label study of the pharmacokinetics and safety of a single dose of moxidectin per oral in subjects aged 4 to 17 years with (or at risk of) onchocerciasis to identify an optimal dose for treatment of children 4 to 11 years
105 MEBENDAZOLE	Efficacy and Safety Of A Single Dose Reigimen And A Multi Dose Regimen Of Mebendazole Against Hookworm Infections in Children And Adolescents In Ghana: A Randomized Control Trail.
400 MEEL 000111 04 71711	
106 MEFLOQCHLOAZITH	A Phase III, Randomized, Opened-Label, Comparative Trial Of Azithromycin Plus Chloroquine Versus Mefloquine For The Treatment Of Uncomplicated Plasmodium Falciparum Malaria In Africa.
MENINGOCOCCAL-A CONJUGATE 107 VACCINE	A Phase II, Double Blind, Randomized, Controlled, Dose Ranging Study to Evaluate the Safety, Immunogenicity Dose Response and Schedule Response of a Meningococcal A Conjugate Vaccine administered concomitantly with local EPI vaccines in Healthy Infants.
MICRONUTRIENT 108 SUPPLEMENTATION	The effect of micronutrient supplementation in combination with healthy lifestyle coaching on nutrition status and well-being: A 6-month intervention study in Ghana. (MICRONUTRIENT SUPPLEMENTATION)
109 MITAPIVAT	A Phase 2/3, Double-Blind, Randomized, Placebo-Controlled, Multicenter Study to Evaluate the Efficacy and Safety of Mitapivat in Subjects With Sickle Cell Disease.
110 MMS	The Use Of A Multiple Micronutrient Supplement In Women Of Reproductive Age
111 MoRiOn	The Efficacy of Rifapentine 900mg/d plus Moxilloxacin 400mg/d given for 14 or 7 days against Onchocerciasis – a Randomized, Controlled, Parallel-Group, Open Label, Phase II Pilot Trial
112 MOSA STUDY	A phase III, multi-country, randomized, placebo-controlled, double-blinded adaptive platform trial to assess the efficacy and safety of treatments for subjects with monkeypox virus disease
113 MOXIDECTIN  MOXIDECTIN-	Randomized, single-ascending dose, Ivermectin-controlled, double-blind, safety, tolerability, pharmacokinetic and efficacy study of orally administered Moxidectin in subjects with Onchocerca volvulus Infection
114 IVERMECTIN	A Phase III Randomized, Single-Ascending-Dose, Ivermectin-Controlled, Double-Blind, Safety, Tolerability, Pharmacokinetic, and Efficacy Study of Orally Administered Moxidectin in Subjects with Onchocerca volvulus Infection:
115 MPZ-MAL 01	A Phase 2a, Multicenter, Open-label, Dose-finding, Dose Escalation Study of Mepiazumab in Adult Patients Diagnosed with Uncomplicated Plasmodium falciparum Malaria
116 MULTIMAL	Multi-Drug Combination-Therapies to prevent the Development of Drug Resistance: Phase II Controlled Clinical Trial Assessing Candidate Regimens of Multiple-Antimalarial Combinations for the Treatment of Uncomplicated Malarial in Africa
MYCOPIROX_LAGR 117 AY	Randomized, open labelled trial to evaluate the efficacy, safety and tolerability of mycopirox vaginal cream in the treatment of mixed infection vaginitis
118 NANOX.ARC	Multicentric study for assessing safety and clinical performance of Nanox.ARC in providing additional information to conventional twodimensional (2D) radiography when evaluating adult individuals with known or suspected radiographic abnormalities
119 NEOSEP 1	An open-label randomized controlled trial comparing novel combination and currently used antibiotic regimens for the empiric treatment of neonatal sepsis with a run-in confirmatory pharmacokinetic phase (NEOSEP 1)
120 NEOVITA	Feasibility Studies
NOGUCHI 121 FILARIASIS	Determination of the Prevalence of LF Infection in Districts Not Included in LF Control Activities and of the Basis for Integrated Implementation of LF - Onchocerciasis Elimination Strategies in Potentially Co-endemic Areas
122 NOGUCHI SCD	A Phase 1B Dose – Finding Pharmacokinetics and Pharmacokynamic Study Oof NVX – 508 In Sickle Cell Disease (SCD) Patients
NON-INVASIVE 123 HAEM DEVICE	A Comparison of Hemoglobin Values as Measured By The Pronto And Pronto 7 Non-Invasive Hemoglobin Devices, The Hemocue Hb 201+, And A Hematology Analyzer Among Pregnant Women Attending Antenatal Care Clinic In Ghana
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124 NOVASIL	Safety and Efficacy Evaluation of Novasil: Strategy for the Protection of Humans from Aflatoxin Toxicity
125 NOVIC TRIAL	Novel vacuum-induced Haemorrhage control for postpartum Haemorrhage: a multicentre randomised trial
126 OXYTOCIN	Determining the Effect of Prophylactic Administration Of Oxytocin In Uniject™ By A Community Health Officer On Post-Partum Haemorrage At Home Births In The Kintampo North And South Districts Of Ghana
127 PEARL	Phase III, randomized, observer-blind, placebo-controlled, multi-center, multinational study to evaluate the efficacy, immunogenicity, and safety of a Respiratory Syncytial Virus vaccine in infants and toddlers (PEARL)
PFCSP_MVACS_MA 128 LARIA	Partial Double-Blind, Randomized Study of PFCSP DNA/MVA Prime Boost Vaccine
129 PIVOT	Prospective Identification of Variables as Outcomes for Treatment (PIVOT): A Phase II clinical trial of hydroxyurea for children and adults with HbSC disease
POLYPHENOL-RICH	
COCOA POWDER 130 TRIAL	Polyphenol-rich Cocoa Powder as Adjuvant Therapy in Patients with Covid-19.
POST	
MASTECTOMY PAIN 131 RELIEF	ULTRASOUND-GUIDED ERECTOR SPINAE PLANE BLOCK FOR POST-MASTECTOMY PAIN RELIEFve
132 PLATINUM	: A multi-part, multi-center PLATform study to assess the efficacy, safety, tolerability and pharmacokinetics of anti-malarial agents administered asmonotherapy and/or combination therapy IN patients with Uncomplicated Plasmodium falciparum Malaria
PMC-RTSS 133 SUBSTUDY	Characterization of the Impact of Combining Perennial Malaria Chemoprevention with RTS,S/AS01E Malaria Vaccination on Vaccine Induced and Naturally Acquired Immunity to Malaria.
134 PMC TRIAL	The impact of a combination of the RTS,S/AS01E malaria vaccine and perennial malaria chemoprevention in Ghanaian children
135 PRAISE	An adaptive, Randomized, Placebo-controlled, Double-Blind, Multi-center Study of Oral FT-4202, a Pyrtuvate Kinase Activator in Patients with Sickle Cell disease (PRAISE)
136 PREGACT	Evaluating the Safety And Efficacy Of Artemisinin-Based Combination Treatments For African Pregnant Women With Malaria
137 PRENABELT	A Maternal Device to Reduce the Risk of Stillbirth and Low-Birth Weight
138 PROBIOTIC	A double-blind randomized control trial of a synbiotic vs. placebo among pregnant women to evaluate colonization of the gut microbiota of their infants with Lactobacillus plantarum (Probiotics pilot in Ghana)
PROBIOTIC(IN MILD COGNITIVE	
139 IMPAIRMENT)	Assessing the Therapeutic Effect of Probiotics on Individuals with Mild Cognitive Impairment
140 PROFUSA	Continuous monitoring of Tissue Oxygen in Septic Patients using an injectable Biosensor
PYRONARIDINE	Committed a first model of Agent in Copie 1 aloch a using an injectable biosensor
ARTESUNATE VRS 141 COARTEM	andomized multicentre clinical study to assess the safety and efficacy of fixed dose formulation of oral pyronaridine artesunate tablet versus coartern in children and adult patients with acute uncomplicated plasmodium falciparium malaria
142 PRCR DIPSTICK	Validation of a Protein Creatinine (PrCr) Dipstick Diagnostic Test for Proteinuria Screening on Antenatal Care Clinics in Ghana
	Validation of a Protein Cleaning (PCC) Disputs Diagnosis Test for Proteining Screening on American Care Clinics in Gillana
143 PRCR SPOT	
	Evaluating the clinical utility and operational fit of the lifeAssay Diagnostics Test-It TM PrCr urinary dipstick test to assess risk of pre- eclampsia in referral hospitals in Ghana: A SPOT nested study, developing and VALidating a Severe Pre-eclampsia adverse Outcome Triage (SPOT) score
144 REALISE	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children
144 REALISE 145 RECOVERY	
145 RECOVERY 146 REVIVE	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children
145 RECOVERY	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children  Randomized Evaluation of Covid-19 Therapy (RECOVERY)
145 RECOVERY  146 REVIVE RIFAMPIN VS	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children  Randomized Evaluation of Covid-19 Therapy (RECOVERY)  Reducing Mortality in Adults with Advanced HIV Disease (REVIVE)
145 RECOVERY  146 REVIVE RIFAMPIN VS 147 ISONIAZID  148 ROBOCOW	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children  Randomized Evaluation of Covid-19 Therapy (RECOVERY)  Reducing Mortality in Adults with Advanced HIV Disease (REVIVE)  A Randomized Clinical Trial of 4 months Rifampin versus 9 months Isoniazid for treating Latent TB Infection  RANDOMIZED PLACEBO-CONTROLLED TRIAL TESTING 0.2% CHLORHEXIDINE MOUTHWASH TO REDUCE POSTOPERATIVE RESPIRATORY TRACT INFECTIONS IN ABDOMINAL SURGERIES
145 RECOVERY  146 REVIVE RIFAMPIN VS 147 ISONIAZID	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children  Randomized Evaluation of Covid-19 Therapy (RECOVERY)  Reducing Mortality in Adults with Advanced HIV Disease (REVIVE)  A Randomized Clinical Trial of 4 months Rifampin versus 9 months Isoniazid for treating Latent TB Infection
145 RECOVERY  146 REVIVE RIFAMPIN VS 147 ISONIAZID  148 ROBOCOW	A Pragmatic Phase III Multi-Centre Clinical Trial to Evaluate the Safety and Effectiveness of a Single Dose of an Albendazole-Ivermectin Coformulation vs Albendazole for Preventive Chemotherapy of Soil-Transmitted Helminth Infections in School-Aged Children  Randomized Evaluation of Covid-19 Therapy (RECOVERY)  Reducing Mortality in Adults with Advanced HIV Disease (REVIVE)  A Randomized Clinical Trial of 4 months Rifampin versus 9 months Isoniazid for treating Latent TB Infection  RANDOMIZED PLACEBO-CONTROLLED TRIAL TESTING 9.2% CHLORHEXIDINE MOUTHWASH TO REDUCE POSTOPERATIVE RESPIRATORY TRACT INFECTIONS IN ABDOMINAL SURGERIES
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