

Pipeline assets and clinical trials appendix Q2 2024

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Opportunity driven



Innovation: Pipeline growth

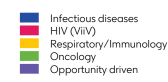
Overview of potential new vaccines and medicines



70 potential new vaccines and medicines in pipeline

Phase III / Registration – 18 assets

Arexvy (RSV vaccine)	Recombinant protein, adjuvanted*	RSV adults (50-59 YoA AIR)^1**
gepotidacin (GSK2140944)	BTI inhibitor*	Uncomplicated UTI**
bepirovirsen (GSK3228836)	Antisense oligonucleotide*	Chronic HBV infection**
Bexsero (MenB vaccine)	Recombinant protein, OMV	Meningitis B (infants US)
MenABCWY vaccine (GSK3536819)	Recombinant protein, OMV, conjugated vaccine	MenABCWY, 1st Gen^
tebipenem pivoxil (GSK3778712)	Antibacterial carbapenem*	Complicated UTI
ibrexafungerp (GSK5458448)	Antifungal glucan synthase inhibitor*	Invasive candidiasis
Nucala (mepolizumab)	Anti-IL5 antibody	COPD
depemokimab (GSK3511294)	Long-acting anti-IL5 antibody*	Asthma**
latozinemab (GSK4527223)	Anti-sortilin antibody*	Frontotemporal dementia ² **
camlipixant (GSK5464714)	P2X3 receptor antagonist	Refractory chronic cough
Low carbon version of MDI ³ , Ventolin (salbutamol)	Beta 2 adrenergic receptor agonist	Asthma
Jemperli (dostarlimab)	Anti-PD-1 antibody*	Endometrial cancer^**
Zejula (niraparib)	PARP inhibitor*	Ovarian cancer**
Blenrep (belantamab mafodotin)	Anti-BCMA ADC*	Multiple myeloma
cobolimab (GSK4069889)	Anti-TIM-3 antibody*	Non-small cell lung cancer
belrestotug (GSK4428859)	Anti-TIGIT antibody*	Non-small cell lung cancer**
linerixibat (GSK2330672)	IBAT inhibitor	Cholestatic pruritus in primary biliary cholangitis





70 potential new vaccines and medicines in pipeline

Phase II – 32 assets

HIV (ViiV)
Respiratory/Immunology
Oncology
Opportunity driven

Infectious diseases

MMRV new strain Shigella Chronic HBV infection¹** MenABCWY, 2nd Gen¹ Varicella new strain Adult pneumococcal disease Paediatric pneumococcal disease Gonorrhoea¹ Seasonal flu COVID-19 Flu H5N1 pre-pandemic¹ Cytomegalovirus¹ Therapeutic herpes simplex virus¹ Chronic HBV infection Invasive non-typhoidal salmonella** Tuberculosis **Tuberculosis Tuberculosis** HIV HIV HIV HIV HIV Systemic sclerosis associated interstitial lung disease Osteoarthritis pain** Atopic dermatitis Alzheimer's disease Pulmonary fibrosis

Asthma²

Cancer

NASH/MASH

Malaria fractional dose

^{*}In-license or other alliance relationship with third party ** Additional indications or candidates also under investigation

70 potential new vaccines and medicines in pipeline

Phase I - 20 assets

GSK3536867	Bivalent conjugate*	Salmonella (typhoid + paratyphoid A)
GSK2556286	Mtb cholesterol dependent inhibitor*	Tuberculosis
GSK3772701	P. falciparum whole cell inhibitor*	Malaria
GSK4024484	P. falciparum whole cell inhibitor*	Malaria
GSK3882347	FimH antagonist*	Uncomplicated UTI
GSK3923868	PI4K beta inhibitor	Rhinovirus disease
GSK3965193	PAPD5/PAPD7 inhibitor	Chronic HBV infection ¹
GSK5251738	TLR8 agonist*	Chronic HBV infection
cabotegravir (GSK1265744)	Integrase inhibitor	HIV
GSK3888130	Anti-IL7 antibody*	Autoimmune disease
GSK3862995	Anti-IL33 antibody	COPD
GSK5462688	RNA-editing oligonucleotide*	Alpha-1 antitrypsin deficiency
GSK4347859	Interferon pathway modulator	Systemic lupus erythematosus
GSK4381562	Anti-PVRIG antibody*	Cancer
XMT-2056 ² (wholly owned by Mersana Theraprutics)	STING agonist ADC*	Cancer
belantamab (GSK2857914)	Anti-BCMA antibody	Multiple myeloma**
GSK4524101	DNA polymerase theta inhibitor*	Cancer ¹
GSK5764227	ADC-targeting B7-H3*	Solid tumors
GSK5733584	ADC-targeting B7-H4*	Gynecologic malignancies
GSK4172239	DNMTI inhibitor*	Sickle cell disease

Infectious diseases HIV (ViiV) Respiratory/Immunology Oncology Opportunity driven



Infectious diseases pipeline

Infectious diseases HIV (ViiV) Respiratory/Immunology Oncology Opportunity driven

Phase III / Registration – 7 assets

Arexvy (RSV vaccine)	Recombinant protein, adjuvanted*	RSV adults (50-59 YoA AIR)^1**
gepotidacin (GSK2140944)	BTI inhibitor*	Uncomplicated UTI**
bepirovirsen (GSK3228836)	Antisense oligonucleotide*	Chronic HBV infection**
Bexsero (MenB vaccine)	Recombinant protein, OMV	Meningitis B (infants US)
MenABCWY vaccine (GSK3536819)	Recombinant protein, OMV, conjugated vaccine	MenABCWY, 1st Gen^
tebipenem pivoxil (GSK3778712)	Antibacterial carbapenem*	Complicated UTI
ibrexafungerp (GSK5458448)	Antifungal glucan synthase inhibitor*	Invasive candidiasis

Phase I - 8 assets

GSK3536867	Bivalent conjugate*	Salmonella (typhoid + paratyphoid A)
GSK2556286	Mtb cholesterol dependent inhibitor*	Tuberculosis
GSK3772701	P. falciparum whole cell inhibitor*	Malaria
GSK4024484	P. falciparum whole cell inhibitor*	Malaria
GSK3882347	FimH antagonist*	Uncomplicated UTI
GSK3923868	PI4K beta inhibitor	Rhinovirus disease
GSK3965193	PAPD5/PAPD7 inhibitor	Chronic HBV infection ²
GSK5251738	TLR8 agonist*	Chronic HBV infection

Phase II - 19 assets

GSK3437949	Recombinant protein, adjuvanted*	Malaria fractional dose
GSK4406371	Live, attenuated	MMRV new strain
GSK3536852	GMMA*	Shigella
GSK3528869	Viral vector with recombinant protein, adjuvanted*	Chronic HBV infection ² **
GSK4023393	Recombinant protein, OMV, conjugated vaccine	MenABCWY, 2 nd Gen ²
GSK4178116	Live, attenuated	Varicella new strain
GSK5101956	MAPS Pneumococcal 24-valent*	Adult pneumococcal disease
GSK5101955	MAPS Pneumococcal 24-valent paed*	Paediatric pneumococcal disease
GSK4348413	GMMA	Gonorrhoea ²
GSK4382276	mRNA*	Seasonal flu
GSK4396687	mRNA*	COVID-19
GSK5536522	mRNA*	Flu H5N1 pre-pandemic ²
GSK3993129	Adjuvanted recombinant subunit	Cytomegalovirus ²
GSK3943104	Recombinant protein, adjuvanted*	Therapeutic herpes simplex virus ²
GSK5637608	Hepatitis B virus-targeted siRNA*	Chronic HBV infection
GSK4077164	Bivalent GMMA*	Invasive non-typhoidal salmonella**
ganfeborole (GSK3036656)	Leucyl t-RNA synthetase inhibitor*	Tuberculosis
sanfetrinem cilexetil (GV118819)	Serine beta lactamase inhibitor*	Tuberculosis
alpibectir (BVL-GSK3729098)	Ethionamide booster*	Tuberculosis



HIV pipeline

Phase II – 5 assets

VH3810109	Broadly neutralizing antibody*	HIV
VH3739937	Maturation inhibitor	HIV
VH4004280	Capsid protein inhibitor	HIV
VH4011499	Capsid protein inhibitor	HIV
VH4524184	Integrase inhibitor*	HIV

Phase I - 1 asset

cabotegravir (GSK1265744) Integrase inhibitor





Respiratory/Immunology pipeline

Phase III / Registration – 5 assets

Nucala (mepolizumab)	Anti-IL5 antibody	COPD
depemokimab (GSK3511294)	Long-acting anti-IL5 antibody*	Asthma**
latozinemab (GSK4527223)	Anti-sortilin antibody*	Frontotemporal dementia ¹ **
camlipixant (GSK5464714)	P2X3 receptor antagonist	Refractory chronic cough
Low carbon version of MDI ² , Ventolin (salbutamol)	Beta 2 adrenergic receptor agonist	Asthma

Phase II – 6 assets

Benlysta (belimumab)	Anti-BLys antibody	Systemic sclerosis associated interstitial lung disease
GSK3858279	Anti-CCL17 antibody*	Osteoarthritis pain**
GSK1070806	Anti-IL18 antibody	Atopic dermatitis
GSK4527226 (AL-101)	Anti-sortilin antibody*	Alzheimer's disease
GSK3915393	TG2 inhibitor*	Pulmonary fibrosis
GSK5784283	TSLP monoclonal antibody*	Asthma ³

Phase I - 4 assets

GSK3888130	Anti-IL7 antibody*	Autoimmune disease
GSK3862995	Anti-IL33 antibody	COPD
GSK5462688	RNA-editing oligonucleotide*	Alpha-1 antitrypsin deficiency
GSK4347859	Interferon pathway modulator	Systemic lupus erythematosus





Oncology pipeline

Phase III / Registration – 5 assets

Jemperli (dostarlimab)	Anti-PD-1 antibody*	Endometrial cancer^**
Zejula (niraparib)	PARP inhibitor*	Ovarian cancer**
Blenrep (belantamab mafodotin)	Anti-BCMA ADC*	Multiple myeloma
cobolimab (GSK4069889)	Anti-TIM-3 antibody*	Non-small cell lung cancer
belrestotug (GSK4428859)	Anti-TIGIT antibody*	Non-small cell lung cancer**

Phase II – 1 asset

nelistotug (GSK6097608) Anti-CD96 antibody*

Phase I – 6 assets

GSK4381562	Anti-PVRIG antibody*	Cancer
XMT-2056 ¹ (wholly owned by Mersana Theraprutics)	STING agonist ADC*	Cancer
belantamab (GSK2857914)	Anti-BCMA antibody	Multiple myeloma**
GSK4524101	DNA polymerase theta inhibitor*	Cancer ²
GSK5764227	ADC-targeting B7-H3*	Solid tumors
GSK5733584	ADC-targeting B7-H4*	Gynecologic malignancies



Infectious diseases HIV (ViiV)

Oncology Opportunity driven

Respiratory/Immunology

Opportunity driven pipeline

Phase III / Registration – 1 asset

linerixibat (GSK2330672) IBAT inhibitor

Cholestatic pruritus in primary biliary cholangitis

Phase II – 1 asset

GSK4532990 HSD17B13 RNA interference*

NASH/MASH

Phase I - 1 asset

GSK4172239

DNMT1 inhibitor*

Sickle cell disease



*In-license or other alliance relationship with third party

Infectious diseases HIV (ViiV)

Oncology Opportunity driven

Respiratory/Immunology

Changes since Q1 2024

Infectious diseases HIV (ViiV) Respiratory/Immunology Oncology Opportunity driven

Changes on pipeline

Progressed from Phase II to Phase III

belrestotug (GSK4428859): Anti-TIGIT antibody, non-small cell lung cancer

New to Phase II

GSK5536522: mRNA, flu H5N1 pre-pandemic

Removed from Registration

Omjjara: JAK1, JAK2 and ACVR1 inhibitor, myelofibrosis¹

Removed from Phase II

GSK4106647: Adjuvanted recombinant protein, adjuvanted, human papillomavirus

Removed from Phase I

GSK3494245: Proteasome inhibitor, visceral leishmaniasis

Achieved pipeline catalysts

Regulatory decisions

Arexvy: Adjuvanted recombinant protein, RSV adults (50-59 YoA AIR²)

Omjjara: JAK1, JAK2 and ACVR1 inhibitor, myelofibrosis

JP

Regulatory submission acceptances

Jemperli³: RUBY (Part 1)⁴, 1L Endometrial cancer EU
Blenrep: DREAMM-7/8, 2L+ Multiple myeloma EU

Late-stage readouts

depemokimab: SWIFT-1/2, asthma — Positive phase III data readout



Upcoming pipeline catalysts: 2024 and 2025



	H2 2024		H1 2025		H2 2025	
Regulatory	Arexvy: 50-59 YoA ¹ AIR ²	EU, JP	gepotidacin: EAGLE-2/3, uUTI ⁷	US	gepotidacin: EAGLE-1, GC ¹³	US
decision	Nucala: CRSwNP ³	JP	MenABCWY vaccine 1st Gen	US	depemokimab: SWIFT-1/2, asthma	US
	Jemperli ⁴ : RUBY (Part 1) ^{5,} 1L EC ⁶	US	Shingrix: 18+ YoA	CN	depemokimab: ANCHOR-1/2, CRSwNP ³	US
			Nucala: CRSwNP ³	CN	Blenrep: DREAMM-7/8, 2L+ MM ⁹	US, EU
			Nucala: MATINEE, COPD ⁸	US	linerixibat: GLISTEN, cholestatic pruritus in PBC ¹²	US
			Blenrep: DREAMM-7/8, 2L+ MM ⁹	JP	_	
			Jemperli ⁴ : RUBY (Part 1) ^{5.} 1L EC ⁶	EU		
Regulatory	gepotidacin: EAGLE-2/3, uUTI ⁷	US	gepotidacin: EAGLE-1, GC ¹³	US	Bexsero (infants US)	US
submission	depemokimab: SWIFT-1/2, asthma	US	depemokimab: SWIFT-1/2, asthma	EU, CN, JP	Arexvy 18-49 YoA ¹ AIR ²	US
acceptance	depemokimab: ANCHOR-1/2, CRSwNP ³	US	depemokimab: ANCHOR-1/2, CRSwNP ³	EU, CN, JP	gepotidacin: EAGLE-J, uUTI ⁷	JP
•	Nucala: MATINEE, COPD ⁸	US	Nucala: MATINEE, COPD ⁸	EU, CN	tebipenem pivoxil: PIVOT-PO, cUTI ¹⁴	US
	Blenrep: DREAMM-7/8, 2L+ MM ⁹	US, JP	Ventolin (low carbon MDI): asthma	EU	camlipixant: CALM-1/2, RCC ¹⁵	US, EU
	Blenrep: DREAMM-7, 2L+ MM ⁹	CN	linerixibat: GLISTEN, cholestatic pruritus in PBC ¹²	US, EU, CN	Blenrep: DREAMM-8, 2L+ MM ⁹	CN
			_		cobolimab ⁴ : COSTAR, 2L NSCLC ¹¹	US, EU
					linerixibat: GLISTEN, cholestatic pruritus in PBC ¹²	JP
Late-stage	depemokimab: ANCHOR-1/2, CRSwNP ³		Arexvy18-49 YoA ¹ AIR ²		Bexsero (infants US)	
Phase III	Nucala: MATINEE, COPD ⁸		Ventolin (low carbon MDI): asthma		tebipenem pivoxil: PIVOT-PO, cUTI ¹⁴	
readouts	Zejula ⁴ : FIRST, 1L maintenance OC ¹⁰		cobolimab⁴: COSTAR, 2L NSCLC ¹¹		camlipixant: CALM-1/2, RCC ¹⁵	
	Zejula ³ : ZEAL, 1L maintenance NSCLC ¹¹				depemokimab: OCEAN, EGPA ¹⁶	
	linerixibat: GLISTEN, cholestatic pruritus in PBC ¹²				depemokimab: NIMBLE, asthma	
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Designations in our pipeline

Infectious diseases HIV (ViiV) Respiratory/Immunology Oncology Opportunity driven

Breakthrough Designation

GSK5101956	MAPS Pneumococcal 24-valent*	Adult pneumococcal disease	
latozinemab (GSK4527223)	Anti-sortilin antibody*	Frontotemporal dementia ²	

Fast Track

gepotidacin (GSK2140944)	BTI inhibitor*	Urogenital gonorrhoea
bepirovirsen (GSK3228836)	Antisense oligonucleotide*	Chronic HBV infection
GSK4382276	mRNA*	Seasonal flu
alpibectir (BVL-GSK3729098)	Ethionamide booster*	Tuberculosis
GSK4348413	GMMA	Gonorrhoea
tebipenem pivoxil (GSK3778712)	Antibacterial carbapenem*	Complicated UTI
ibrexafungerp (GSK5458448)	Antifungal glucan synthase inhibitor*	Invasive candidiasis
GSK3858279	Anti-CCL17 antibody*	Osteoarthritis pain
GSK3858279	Anti-CCL17 antibody*	Diabetic peripheral neuropathic pain
latozinemab (GSK4527223)	Anti-sortilin antibody*	Frontotemporal dementia ²
Jemperli ¹ (dostarlimab)	Anti-PD-1 antibody*	Neoadjuvant dMMR/MSI-H 1L rectal cancer
GSK4172239	DNMT1 inhibitor*	Sickle cell disease

Priority Review

Jemperli ¹ (dostarlimab) Anti-P[D-1 antibody*	Endometrial cancer^

Orphan Drug Designation

	_	
ibrexafungerp (GSK5458448) US, EU	Antifungal glucan synthase inhibitor*	Invasive candidiasis
Benlysta (belimumab) US	Anti-BLys antibody	Systemic sclerosis associated interstitial lung disease
latozinemab (GSK4527223) US, EU	Anti-sortilin antibody*	Frontotemporal dementia ²
depemokimab (GSK3511294) JP	Long-acting anti-IL5 antibody*	Hypereosinophilic syndrome
linerixibat (GSK2330672) US, EU	IBAT inhibitor	Cholestatic pruritus in primary biliary cholangitis

Qualified Infectious Disease Product Designation

gepotidacin (GSK2140944)	BTI inhibitor*	Uncomplicated UTI
gepotidacin (GSK2140944)	BTI inhibitor*	Urogenital gonorrhoea
ebipenem pivoxil (GSK3778712)	Antibacterial carbapenem*	Complicated UTI
brexafungerp (GSK5458448)	Antifungal glucan synthase inhibitor*	Invasive candidiasis

BREAKTHROUGH DESIGNATION (US) – a process designed to expedite the development and review of medicines intended to treat serious conditions, where preliminary clinical evidence indicates the drug may demonstrate substantial improvement over available therapy

FAST TRACK (US) - a program designed to facilitate the expedited development and review of medicines to treat serious conditions and fill an unmet medical need

PRIORITY REVIEW (US) – indicated the US FDA's goal to take action on an application within 6 months (compared to 10 months under standard review)

OPHAN DRUG DESIGNATION – intended for treatment, diagnosis or prevention of rare disease/disorders that affect fewer than 200,000 patients in the US, or not more than 5 in 10,000 in the EU or that affect more than this number of patients but are not expected to recover the costs of developing and marketing a treatment drug, or if intended for use in less than 50,000 patients in Japan and for which there is a high medical need

QUALIFIED INFECTIOUS DISEASE PRODUCT DESIGNATION (US) - an antibacterial or antifungal drug for human use intended to treat serious or lifethreatening infections



*In-license or other alliance relationship with third party ^ In registration 1. Tesaro asset 2. In patients with progranulin gene mutation

Clinical Trials





Phase

Infectious diseases

Arexvy (RSV Adults)

NCT04732871 - RSV OA=ADJ-004

Phase	III
Patient	Adults ≥60 years of age
Subjects	1720
	Arm A: RSVPreF3 OA Day 1, 12 months & 24 months
Treatment arms	Arm B: RSVPreF3 OA Day 1, 24 and 48 months
	Arm C: RSVPreF3 OA Day 1 then follow up, at month 36, re-randomization in 2 groups
Description	A randomised, open-label, multi-country trial to evaluate the immunogenicity, safety, reactogenicity and persistence of a single dose of the RSVPreF3 OA investigational vaccine and different revaccination schedules in adults aged 60 years and above
Timeline	Trial start: Q1 2021
Ilmeline	Primary data reported: Q2 2022
Key end points	Humoral immune response
Clinicaltrials.gov	<u>Link</u>

NCT04886596 - RSV OA=ADJ-006

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Patient	Adults ≥60 years of age
Subjects	26,668
	Arm A: RSVPreF3 OA Lot 1
	Arm B: RSVPreF3 OA Lot 2
Treatment arms	Arm C: RSVPreF3 OA Lot 3
	Arm D: RSVPreF3 OA Lot 4
	Arm E: Placebo
Description	A randomised, placebo-controlled, observer-blind, multi-country trial to demonstrate the efficacy of a single dose and revaccination prior to Season 2 of GSK's RSVPreF3 OA investigational vaccine in adults aged 60 years and above
T: !:	Trial start: Q2 2021
Timeline	Primary data reported: Q2 2022; season two data reported Q2 2023
Key end points	Efficacy of a single dose and revaccination prior to Season 2 of RSVPreF3 OA vaccine in the prevention of RSV-LRTD in adults ≥ 60 yoa
Clinicaltrials.gov	<u>Link</u>



Arexvy (RSV Adults)

NCT04841577 - RSV OA=ADJ-007

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Phase	III	Phase	III
Patient	Adults ≥60 years of age	Patient	Adults aged 65 years and above
Subjects	976	Subjects	1029
T	Arm A: 1 dose of RSVPreF3 OA +1 dose of FLU-QIV on Day 1	Tue seture and summer	Arm A: 1 dose of RSVPreF3 OA + 1 dose of Flu-HD on day 1
Treatment arms	Arm B: 1 dose of FLU-QIV on Day 1, 1 dose of RSVPreF3 OA on Day 31	Treatment arms	Arm B: 1 dose of Flu HD on Day 1,1 dose of RSVPreF3 OA on Day 31
Description	An open-label, randomised, controlled, multi-country trial to evaluate the immune response, safety and reactogenicity of RSVPreF3 OA investigational vaccine when co-administered with FLU-QIV vaccine in adults aged 60 years and above	Description	An open-label, randomised, controlled, multi-country trial to evaluate the immune response, safety and reactogenicity of RSVPreF3 OA investigational vaccine when co-administered with FLU HD vaccine in adults aged 65 years and above
Time elim e	Trial start: Q2 2021	Timeline	Trial start: Q4 2022
Timeline	Primary data reported: Q4 2022	Timeline	Primary data reported: Q2 2023
Key end points	Humoral immune response 1 month post vaccination upon co-administration compared to the immune response when vaccine is administered alone	Key end points	Humoral immune response 1 month post vaccination upon co-administration compared to the immune response when vaccine is administered alone
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>



Arexvy (RSV Adults)

NCT05059301 - RSV OA=ADJ-009

Phase	III
Patient	Adults aged 60 years and above
Subjects	770
	Arm A: 1 dose of a combination of the RSVPreF3 antigen Lot 1 and AS01E adjuvant Lot A at day 1
Treatment arms	Arm B: 1 dose of a combination of the RSVPreF3 antigen Lot 2 and AS01E adjuvant Lot B at day 1
	Arm C: 1 dose of a combination of the RSVPreF3 antigen Lot 3 and AS01E adjuvant Lot C at Day 1
Description	A randomised, double-blind, multi-country trial to evaluate consistency, safety and reactogenicity of 3 lots of RSVPreF3 OA investigational vaccine administrated as a single dose in adults aged 60 years and above
Timeline	Trial start: Q4 2021
I imeline	Trial end: Q2 2022
Key end points	RSVPreF3-binding IgG concentrations at 1 month post vaccination for three lots of RSVPreF3 OA investigational vaccine
Clinicaltrials.gov	<u>Link</u>

NCT05568797 - RSV OA=ADJ-017

Phase	III
Patient	Adults aged 65 years and above
Subjects	1045
Treatment arms	Arm A: 1 dose RSVPreF3 OA investigational vaccine and 1 dose of FLU aQIV vaccine on Day 1
	Arm B: one dose of Flu aQIV on day 1 and 1 dose of RSVPreF3 OA on day 31
Description	An open-label, randomised, controlled, multi-country trial to evaluate the immune response, safety and reactogenicity of an RSVPreF3 OA investigational vaccine when co-administered with FLU aQIV (inactivated influenza vaccine – adjuvanted) in adults aged 65 years and above
Timeline	Trial start: Q4 2022
Ilmeline	Primary data reported: Q2 2023
Key end points	Humoral immune response 1 month post vaccination upon co-administration compared to the immune response when vaccine is administered alone
Clinicaltrials.gov	<u>Link</u>



Innovation: Pipeline growth Infectious diseases HIV Respiratory/Immunology Oncology Opportunity driven Glossary

Infectious diseases

Arexvy (RSV Adults)

NCT05590403 - RSV OA-018

Phase	III
Patient	Adults 50-59 years of age, including adults at increased risk of respiratory syncytial virus lower respiratory tract disease, and older adults ≥60 years of age
Subjects	1576
	Arm A: adults HA-RSVPreF3 OA Group
	Arm B: adults HA-Placebo Group
Treatment arms	Arm C: adults AIR-RSVPreF3 OA Group
	Arm D: adults AIR-Placebo Group
	Arm E: OA-RSVPReF3 OA Group ≥60 years of age
Description	An observer-blind, randomised, placebo-controlled trial to evaluate the non-inferiority of the immune response and safety of the RSVPreF3 OA investigational vaccine in adults 50 59 years of age, including adults at increased risk of respiratory syncytial virus lower respiratory tract disease, compared to older adults ≥60 years of age
T :	Trial start: Q4 2022
Timeline	Primary data reported: Q4 2023
Key end points	Humoral immune response in healthy participants 50-59 years of age and in participants 50-59 years of age at increased risk of RSV-LRTD compared to OA (≥ 60 yoa)
Clinicaltrials.gov	<u>Link</u>

NCT05879107 - RSV OA=ADJ-019

Phase	III
Patient	Adults ≥60 years of age
Subjects	1113
Treatment arms	Arm A (co-ad group): RSVPreF3 OA investigational vaccine co-administered with PCV20 vaccine
	Arm B (control group): PCV20 vaccine on Day 1 and the RSVPreF3 OA investigational vaccine on Day 31.
Description	An open-label, randomised, controlled, multi-country study to evaluate the immune response, safety and reactogenicity of RSVPreF3 OA investigational vaccine when co-administered with PCV20 in adults aged 60 years and older
Timeline	Trial start: Q2 2023
Timeline	Data anticipated: H2 2024
Key end points	Opsonophagocytic antibody titers for each of the pneumococcal vaccine serotypes and RSV-A & RSV-B serum neutralizing titers
Clinicaltrials.gov	<u>Link</u>

Arexvy (RSV Adults)

NCT05966090 - RSV OA=ADJ-020

NCT05921903 - RSV OA=ADJ-	023

Phase	III	Phase	IIb
Patient	Adults aged 50 years and older	Patient	Immunocompromised (IC) adults 50 years of age and above
Subjects	530	Subjects	375
	Arm A: Participants will be administered first dose of HZ/su vaccine and the RSVPreF3 OA investigational vaccine together on Day 1. A second dose of the		Arm A: RSV_IC_1 group, IC patients receiving 1 dose of RSVPreF3 OA investigational vaccine at Visit 1 (Day 1).
Treatment arms	HZ/su vaccine will be administered at Day 61. Arm B: Participants will be administered first dose HZ/su vaccine on Day 1,	Treatment arms	Arm B: RSV_IC_2 group, IC patients receiving 2 doses of RSVPreF3 OA investigational vaccine at Visit 1 (Day 1) and Visit 3 (Visit 1 + 30-60 days)
	followed by the RSVPreF3 OA investigational vaccine on Day 31, and then second dose of HZ/su vaccine on Day 61.		Arm C: RSV_HA group, healthy participants receiving 1 dose of RSVPreF3 OA investigational vaccine at Visit 1 (Day 1).
Description	An open-label, randomised, controlled, multi-country study to evaluate the immune response, safety and reactogenicity of RSVPreF3 OA investigational vaccine when co-administered with Herpes Zoster recombinant subunit (HZ/su) vaccine in adults aged 50 years and older	Description	A randomised, controlled, open-label trial to evaluate the immune response and safety of the RSVPreF3 OA investigational vaccine in adults (≥50 years of age) when administered to lung and renal transplant recipients comparing one versus two doses and compared to healthy controls (≥50 years of age)
T. I.	Trial start: Q3 2023		receiving one dose
Timeline	Data anticipated: H2 2024	T'	Trial start: Q3 2023
Key end points	Anti-gE antibody concentrations expressed as group geometric mean	Timeline	Data anticipated: H2 2024
	concentration ratio RSV-A & -B serum neutralizing titers expressed as group geometric mean titer	Key end points	RSV-A & -B serum neutralizing titers expressed as mean geometric increase post Dose 2 over post Dose 1
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>



Arexvy (RSV Adults)

NCT06374394 - RSV OA=ADJ-013

Phase	III
Patient	Adults aged 50 years and above
Subjects	850
Treatment arms	RSVPreF3 OA investigational vaccine
	COVID-19 mRNA vaccine
Description	An open-label, randomized, controlled study to evaluate the immune response, safety and reactogenicity of RSVPreF3 OA investigational vaccine when co-administered with a COVID-19 mRNA vaccine (Omicron XBB.1.5)
Timeline	Trial start: Q2 2024
	Data anticipated: H2 2024
Key end points	RSV-A, RSV-B neutralization titers
	SARS-CoV-2 Omicron XBB.1.5 neutralization titers
Clinicaltrials.gov	<u>Link</u>

NCT06389487 - RSV OA=ADJ-025

Phase	IIIb
Patient	Adult participants, 18-49 YOA, at increased risk (AIR) for RSV disease and older adults (OA) participants, >=60 YOA
Subjects	1450
	Part A: RSV-A-AIR Group, RSVPreF3 OA investigational vaccine
Treatment arms	Part A: RSV-OA Group, RSVPreF3 OA investigational vaccine
	Part B: RSV-A-AIR Group, RSVPreF3 OA investigational vaccine
Description	An open-label study to evaluate the non-inferiority of the immune response and to evaluate the safety of the RSVPreF3 OA investigational vaccine in adults 18-49 years of age at increased risk for Respiratory Syncytial Virus disease, compared to older adults >=60 years of age
T:	Trial start: Q2 2024
Timeline	Data anticipated: H2 2024
V	RSV-A, RSV-B neutralizing titers
Key end points	Seroresponse rate (SRR) in RSV-A and RSV-B neutralizing titers
Clinicaltrials.gov	<u>Link</u>



Infectious diseases gepotidacin

NCT04010539 - EAGLE 1

Phase	III
Patient	Uncomplicated urogenital gonorrhoea caused by Neisseria gonorrhoeae
Subjects	628
Treatment arms	Arm A: 2 x 3000 mg gepotidacin for one day
	Arm B: ceftriaxone (500mg IM), 1 g azithromycin
Description	A randomised, multicentre, open-label trial in adolescent and adult participants comparing the efficacy and safety of gepotidacin to ceftriaxone plus azithromycin in the treatment of uncomplicated urogenital gonorrhoea caused by <i>Neisseria gonorrhoeae</i>
Timeline	Trial start: Q4 2019
	Data reported: Q1 2024
Key end points	Number of participants with culture-confirmed bacterial eradication 4-8 days post treatment
Clinicaltrials.gov	<u>Link</u>



Infectious diseases gepotidacin

NCT04020341 - EAGLE 2

Phase	III
Patient	Females with uUTI / acute cystitis
Subjects	1531
Treatment arms	Arm A: 1500 mg BID gepotidiacin + placebo x 5 days
	Arm B: 100 mg BID nitrofurantoin + placebo x 5 days
Description	A randomised, multicentre, parallel-group, double-blind, double-dummy trial in adolescent and adult female participants comparing the efficacy and safety of gepotidacin to nitrofurantoin in the treatment of uncomplicated urinary tract infection (acute cystitis)
Time aline	Trial start: Q4 2019
Timeline	Data reported: Q2 2023
Key end points	Number of participants with therapeutic response (combined per participant clinical and microbiological response)
Clinicaltrials.gov	<u>Link</u>

NCT04187144 - EAGLE 3

Phase	III
Patient	Females with uUTI / acute cystitis
Subjects	1606
Tue orthogonal comme	Arm A: 1500 mg BID gepotidiacin + placebo x 5 days
Treatment arms	Arm B: 100 mg BID nitrofurantoin + placebo x 5 days
Description	A randomised, multicentre, parallel-group, double-blind, double-dummy trial in adolescent and adult female participants comparing the efficacy and safety of gepotidacin to nitrofurantoin in the treatment of uncomplicated urinary tract infection (acute cystitis)
T: I:	Trial start: Q2 2020
Timeline	Data reported: Q2 2023
Key end points	Number of participants with therapeutic response (combined per participant clinical and microbiological response)
Clinicaltrials.gov	<u>Link</u>



bepirovirsen

NCT05630807 - B-WELL 1

Phase	III
Patient	Non-cirrhotic nucleos(t)ide analogue treated patients with chronic hepatitis B virus
Subjects	941
Treatment arms	Arm A: bepirovirsen for 24 weeks
	Arm B: placebo
Description	A multicentre, randomised, double blind trial to confirm the efficacy and safety of treatment with bepirovirsen in participants with chronic hepatitis B virus
Timeline	Trial start: Q1 2023
	Data anticipated: 2026+
Key end points	Number of participants with baseline HBsAg≤ 3000IU/mL achieving functional cure (FC)
Clinicaltrials.gov	<u>Link</u>

NCT05630820 - B-WELL 2

Phase	III
Patient	Non-cirrhotic nucleos(t)ide analogue treated patients with chronic hepatitis B virus
Subjects	859
Treatment arms	Arm A: bepirovirsen for 24 weeks
reatment arms	Arm B: placebo
Description	A multicentre, randomised, double blind trial to confirm the efficacy and safety of treatment with bepirovirsen in participants with chronic hepatitis B virus
Time aline	Trial start: Q1 2023
Timeline	Data anticipated: 2026+
Key end points	Number of participants with baseline HBsAg≤ 3000IU/mL achieving functional cure (FC)
Clinicaltrials.gov	<u>Link</u>



bepirovirsen

Phase	II .
Patient	HBV suppressed subjects under nucleo(s)tide treatment
Subjects	184
Treatment arms	ASO24-targeted immunotherapy group (GSK3228836 (24-week treatment) followed by GSK3528869A)
	ASO24 group (GSK3228836 (24-week treatment) followed by non-active control)
	ASO12-targeted immunotherapy group (GSK3228836 (12-week treatment) followed by GSK3528869A)
	ASO12 group (GSK3228836 (12-week treatment) followed by non-active control)
Description	A single-blinded, randomised, controlled multi-country trial to evaluate the safety, reactogenicity, efficacy and immune response following sequential treatment with an anti-sense oligonucleotide against Chronic Hepatitis B (CHB) followed by Chronic Hepatitis B Targeted Immunotherapy (CHB-TI) in CHB patients receiving nucleos(t)ide analogue (NA) therapy
Timeline	Trial start: Q2 2022 Data anticipated: 2026+
Key end points	Number of subjects reporting local and general AEs and percentage of participants with sustained virologic response
Clinicaltrials.gov	<u>Link</u>



MenABCWY

NCT04707391 - MenABCWY-019

Phase	IIIb	
Patient	Healthy adolescents and adults aged 15-25 years	
Subjects	1247	
	Arm A: 2 doses of MenABCWY days 1, 181 + placebo day 211	
Treatment arms	Arm B: 1 dose MenABCWY day 1; 2 doses of MenB on Day 181 and Day 211	
Description A randomised, controlled, observer-blind trial to evaluate safety and immunogenicity of GSK's meningococcal ABCWY vaccine when administ in healthy adolescents and adults previously primed with meningococcal ACWY vaccine		
Timeline	Trial start: Q1 2021	
i imeline	Data reported: Q1 2024	
Key end points	hSBA titers	
Clinicaltrials.gov	<u>Link</u>	

NCT04502693 - MenABCWY V72 72

Phase	III		
Patient	Healthy adolescents and adults ages 10-25 years		
Subjects	3638		
	Arm A: rMenB+OMV NZ (2/3 dose schedule) plus MenACWY		
	Arm B: rMenB+OMV NZ (2 dose schedule) plus MenACWY plus placebo		
Treatment arms	Arm C: placebo + MenABCWY lot 1		
reatment arms	Arm D: placebo + MenABCWY lot 2		
	Arm E: placebo + MenABCWY lot 3		
	Arm F: rMenB+OMV NZ + MenACWY + placebo		
Description	Effectiveness of GSK Biologicals S.A.'s Meningococcal Group B and combined ABCWY vaccines in healthy adolescents and young adults		
Time alim a	Trial start: Q3 2020		
Timeline	Data reported: Q1 2023		
Key end points	hSBA titers		
Clinicaltrials.gov	, <u>Link</u>		



MenABCWY

NCT05087056 - MenABCWY-020

Patient Healthy adolescents ≥11 to <15 years of age Subjects 300 Treatment arms Arm A: ABCWY-24 Group Arm B: ABCWY-48 Group	Phase	Ilb	
Arm A: ABCWY-24 Group Treatment arms	Patient	Healthy adolescents ≥11 to <15 years of age	
Treatment arms	Subjects	300	
	T	Arm A: ABCWY-24 Group	
•	i reatment arms	Arm B: ABCWY-48 Group	
A randomised, observer-blind trial to describe the safety, tolerability and immunogenicity of MenABCWY administered on different dosing schedules in healthy adolescents	Description	immunogenicity of MenABCWY administered on different dosing schedules in	
Trial start: Q4 2021	T. I.	Trial start: Q4 2021	
Timeline Data anticipated: 2026+	Imeline	Data anticipated: 2026+	
Key end points hSBA titers ≥ LLOQ of each <i>N. meningitidis</i> serogroup B indicator strain	Key end points	hSBA titers ≥ LLOQ of each <i>N. meningitidis</i> serogroup B indicator strain	
Clinicaltrials.gov <u>Link</u>	Clinicaltrials.gov	<u>Link</u>	



GSK3437949 (Malaria fractional dose)

Phase	IIb	
Patient	Children aged 5-17 months	
Subjects	1500	
Treatment arms	R012-20 Group: a full dose of RTS,S/AS01E at Month 0, Month 1, Month 2 and Month 20 R012-14-mD Group: a full dose of RTS,S/AS01E at Month 0, Month 1, Month 2 Month 14, Month 26, Month 38 Fx012-14-mFxD Group: a full dose of RTS,S/AS01E at Month 0, Month 1 and RTS,S/AS01E 1/5th dose at Month 2, Month 14, Month 26, Month 38 Fx017-mFxD Group: a full dose of RTS,S/AS01E at Month 0, Month 1 and RTS,S/AS01E 1/5th dose at Month 7, Month 20, Month 32 Control Group: Subjects will receive rabies vaccine at Month 0, Month 1, Month 2	
Description	A randomized, open-label, controlled, multi-centre trial of the efficacy, safety and immunogenicity of GSK Biologicals' candidate malaria vaccine RTS,S/AS01E evaluating schedules with or without fractional doses, early Dose 4 and yearly doses, in children 5-17 months of age living in sub-Saharan Africa.	
Timeline	Trial start: Q3 2017 Data publication: H2 2024	
Key end points	Incremental efficacy of a schedule with a fractional third dose at Month 2 over the standard schedule. To demonstrate the superiority of a 3-dose schedule of GSK Biologicals' malaria vaccine RTS,S/AS01E with a fractional third dose at Month 2 compared to a standard schedule of RTS,S/AS01E with three full doses in terms of vaccine efficacy against clinical malaria (primary case definition) over 12 months post-Dose 3.	
Clinicaltrials.gov	<u>Link</u>	



GSK4406371 (MMRV new strain vaccine)

Phase	II	
Patient	Healthy children 4-6 years of age	
Subjects	800	
	Investigational MMRV(H)NS vaccine	
Treatment arms	Investigational MM(H)RVNS vaccine	
reatment arms	Investigational M(L)M(L)R(L)V(L)NS vaccine	
	Marketed MMRV_Lot 1 and Lot 2 vaccine	
Description	A single-blind, randomized, controlled trial to evaluate the immunogenicity and safety of a measles, mumps, rubella, varicella vaccine compared with ProQuad, administered in healthy children 4-6 years of age	
Timeline	Trial start: Q4 2022	
i imeline	Data anticipated: H2 2024	
Key end points Anti-measles, anti-mumps, anti-rubella, and anti-glycoprotein H antibogeometric mean concentrations		
Clinicaltrials.gov	<u>Link</u>	



GSK3536852 (Shigella)

Phase	I/II	
Patient	Adults in Europe (Stage 1) followed by age de-escalation from adults to children and infants and dose finding in infants in Africa (Stage 2)	
Subjects	550	
	Drug: altSonflex Placebo (adults stage 1 in Europe)	
	Biological: altSonflex1-2-3 High Dose C (adults stage 1 in Europe, adults, children and infants stage 2 in Africa)	
	Biological: altSonflex1-2-3 Medium Dose B (children and infants stage 2 in Africa)	
Treatment arms	Biological: altSonflex1-2-3 Low Dose A (infants stage 2 in Africa)	
	Comparators: Menveo and Boostrix (adults stage 2 in Africa)	
	Comparators: Menveo and Typhim Vi (children stage 2 in Africa)	
	Comparators: Menveo and Infanrix (infants stage 2 in Africa)	
Description	A staged observer-blind, randomised, controlled, multi-country trial to evaluate the safety, reactogenicity, and immune responses to the GVGH altSonflex1-2-3 vaccine against <i>S. sonnei</i> and <i>S. flexneri</i> serotypes 1b, 2a, and 3a, in adults in Europe (Stage 1) followed by age de-escalation from adults to children and infants, and dose-finding in infants in Africa (Stage 2)	
T. I.	Trial start: Q4 2021	
Timeline	Data anticipated: 2025	
Key end points	Immune response to identify the preferred dose of each component of the altSonflex1-2-3 vaccine (low, medium, or high) for infants 9 months of age in Africa (Stage 2). To evaluate the safety and reactogenicity of the altSonflex1-2-3 vaccine in all participants in Europe and Africa (Stage 1 and Stage 2)	
Clinicaltrials.gov	<u>Link</u>	



GSK3528869 (Chronic HBV infection)

Phase	1/11	
Patient	HBV suppressed subjects under nucleo(s)tide treatment	
Subjects	148	
	ChAd155-hli-HBV low dose formulation	MVA-HBV low dose formulation
Treatment arms	ChAd155-hIi-HBV high dose formulation	MVA-HBV high dose formulation
i reatment arms	HBc-HBs/AS01B-4 low dose formulation	Placebo
	HBc-HBs/AS01B-4 high dose formulation	
Description	A first time in human trial on GSK's therapeutic vaccines to evaluate the reactogenicity, safety, immunogenicity and efficacy on reduction of serum HBV surface antigen in HBV suppressed subjects under nucleo(s)tide treatment.	
Time aline	Trial start: Q1 2019	
Timeline	Data anticipated: 2025	
Key end points	Safety and reactogenicity, as well as percentage of patients with >1 log decline of HBsAg	
Clinicaltrials.gov	<u>Link</u>	



GSK4023393 (MenABCWY, 2nd Gen)

NCT04886154

Phase	1/11	
Patient	Healthy adults (phase I) and healthy adolescents and adults (phase II)	
Subjects	1439	
	Combination Product: MenABCWY-2Gen low dose vaccine	
	Combination Product: MenABCWY-2Gen high dose vaccine	
Treatment arms	Combination Product: Placebo	
	Combination Product: MenB vaccine	
	Biological: MenACWY vaccine	
Description	A randomised, controlled trial to assess the safety, effectiveness and immune response of meningococcal combined ABCWY vaccine when administered to healthy adults (phase I) and to healthy adolescents and adults (phase II)	
	Trial start: Q2 2021	
Timeline	Data anticipated: H1 2024	
Key end points	AEs, including all SAEs, AEs leading to withdrawal and AEs of special interest (AESIs)	
	Immunological vaccine effectiveness by enc-hSBA and immunogenicity by hSBA on indicator strains	
Clinicaltrials.gov	v <u>Link</u>	

NCT05082285

Phase

THOSE	"	
Patient	Healthy infants	
Subjects	724	
	Combination Product: MenABCWY-2Gen low dose vaccine	
Treatment arms	Combination Product: MenABCWY-2Gen high dose vaccine	
i reatment arms	Combination Product: MenABCWY	
	Combination Product: MenB + MenACWY-TT	
Description	A randomised, partially blinded trial to assess the safety, tolerability and immunogenicity of meningococcal combined ABCWY vaccine when administered to healthy infants	
Timeline	Trial start: Q4 2021	
	Data anticipated: 2025	
Key end points	AEs, including all SAEs, AEs leading to withdrawal and AEs of special interest (AESIs), medical attended events (MAE)	
	Immunogenicity by hSBA to indicator strains	
Clinicaltrials.gov	<u>Link</u>	



GSK4178116 (Varicella new strain)

Phase	II .	
Patient	Healthy children between 12-15 months	
Subjects	800	
	Arm A: low potency varicella NS vaccine, plus routine schedule	
	Arm B: medium potency varicella NS vaccine, plus routine schedule	
Treatment arms	Arm C: high potency varicella NS vaccine, plus routine schedule	
	Arm D: marketed varicella vaccine lot 1, plus routine schedule	
	Arm E: marketed varicella vaccine lot 2, plus routine schedule	
Description	A observer-blind, randomised, controlled trial to evaluate the immunogenicity and safety of a varicella vaccine at various potencies compared with Varivax as a first dose, administered in healthy children in their second year of life	
Timeline	Trial start: Q4 2021	
	Data anticipated: H1 2024	
Key end points	Anti-glycoprotein-E antibodies at day 43	
Clinicaltrials.gov	<u>Link</u>	



GSK5101955 (Paediatric Pneumococcal disease)

Phase	II	
Patient	Healthy infants	
Subjects	121	
	Arm A: 1 mcg AFX3772 administered intramuscularly 4 times within 12 months	
Treatment arms	Arm B: 2 mcg AFX3772 administered intramuscularly 4 times within 12 months	
rreatment arms	Arm C: 5 mcg AFX3772 administered intramuscularly 4 times within 12 months	
	Arm D: PCV13 administered intramuscularly 4 times within 12 months	
Description	A randomised, double-blind, multi-dose, dose finding trial to evaluate the safety, tolerability and immunogenicity of AFX3772 compared with PCV13 in healthy infants	
The all a	Trial start: Q2 2022	
Timeline	Data anticipated: 2026+	
Key end points Safety, tolerability profiles of 3 different dose levels of AFX3772 comparation points PCV13 with respect to the proportion of participants with AEs		
Clinicaltrials.gov	<u>Link</u>	



GSK4348413 (Gonorrhoea)

Phase	1/11		
Patient	Healthy adults 18 to 50 years of age		
Subjects	1004		
	Phase I	Phase II	
	NgG low dose investigational vaccine	NgG HTD investigational vaccine	
Treatment arms	NgG medium dose investigational vaccine	NgG below HTD investigational vaccine	
	NgG high dose investigational vaccine	Placebo	
	Placebo		
Description	An observer-blind, randomized, placebo-controlled multi-country trial to assess safety and efficacy of GSK <i>Neisseria</i> gonorrhoeae GMMA (NgG) investigational vaccine when administered to healthy adults 18 to 50 years of age		
Time aline	Trial start: Q4 2022		
Timeline	Data anticipated: 2025		
Key end points	AEs and SAEs		
	Incidence rates of gonorrhoeae in trial phase II		
Clinicaltrials.gov	<u>Link</u>		



GSK4382276 (mRNA Seasonal Flu)

NCT05446740

Phase	T .	
Patient	Healthy younger and older adults	
Subjects	324	
	GSK4382276A Dose level 1	GSK4382276A Dose level 8
	GSK4382276A Dose level 2	GSK4382276A Dose level 9
T	GSK4382276A Dose level 3	GSK4382276A Dose level 10
Treatment arms	GSK4382276A Dose level 4	GSK4382276A Dose level 11
	GSK4382276A Dose level 6	Combination Product: FDQ21A-NH
	GSK4382276A Dose level 7	Combination Product: FDQ22A-NH
Description	A randomized, observer-blind, dose-escalation trial to evaluate the safety, reactogenicity and immunogenicity of an mRNA-based monovalent influenza vaccine candidate in healthy younger and older adults	
Time aline	Trial start: Q3 2022	
Timeline	Final data anticipated: H1 2024	
Key end points	Safety and reactogenicity, including number of participants reporting systemic and solicited administration site events	
	Serum anti-influenza seroconversion ro	ates and geometric mean titers
Clinicaltrials.gov	<u>Link</u>	

NCT05823974

Phase

1/11

Patient	Healthy younger and older adults
Subjects	1256
	Biological: Flu mRNA
Treatment arms	Combination Product: Control 1
	Combination Product: Control 2
Description	A randomized, dose-finding/dose-confirmation study to evaluate the reactogenicity, safety and immunogenicity of mRNA-based multivalent seasonal influenza vaccine candidates administered in healthy younger and older adults
Time elim e	Trial start: Q2 2023
Timeline	Final data anticipated: H2 2024
Key end points	Safety and reactogenicity, including number of participants reporting systemic and solicited administration site events
	Serum anti-influenza antigen seroconversion rates and geometric mean titers
Clinicaltrials.gov	<u>Link</u>



GSK4396687 (mRNA COVID-19)

Phase	II
Patient	Adults at least 18 years old
Subjects	675
	Arm A: CV0701 bivalent high dose
	Arm B: CV0701 bivalent medium dose
Treatment arms	Arm C: CV0701 bivalent low dose
reatment arms	Arm D: CV0601 monovalent high dose
	Arm E: Control vaccine
	Arm F: CV0801 Monovalent
Description	A randomized, active-controlled, observer-blind study to assess the safety, reactogenicity, and immunogenicity of a booster dose of investigational COVID-19 mRNA vaccines in healthy adults who previously received a complete primary vaccination series with or without booster dose(s)
T' !'	Trial start: Q3 2023
Timeline	Data anticipated: H2 2024
Key end points	Serum neutralizing titers against pseudoviruses bearing SARS-CoV-2 spike proteins at Day 29
	Percentage of participants with solicited local AE during 7 days after vaccination
Clinicaltrials.gov	<u>Link</u>



GSK5536522 (mRNA Flu H5N1 pre-pandemic)

Phase	1/11
Patient	Healthy younger and older adults
Subjects	1080
Treatment arms	Phase 1 cohort 1: Flu Pandemic mRNA (5 dose levels) and placebo Phase 1 cohort 2: Flu Pandemic mRNA (5 dose levels) and placebo Phase 2 Part A cohort 3: Flu Pandemic mRNA (5 dose levels) or placebo Phase 2 Part A cohort 4: Flu Pandemic mRNA (5 dose levels) or placebo Phase 2 Part B cohort 5: Flu Pandemic mRNA (7 dose levels) or placebo Phase 2 Part B cohort 6: Flu Pandemic mRNA (7 dose levels) or placebo
Description	A randomized, observer-blind, dose-finding/dose-confirmation study to evaluate the safety, reactogenicity and immunogenicity of the mRNA-based investigational pandemic H5 influenza vaccine candidate administered in healthy younger and older adults
Timeline	Trial start: Q2 2024 Data anticipated: 2025
Key end points	Percentage of participants with AEs, MAAEs, SAEs, and AESIs.
Clinicaltrials.gov	<u>Link</u>



GSK3993129 (CMV)

Phase	1/11
Patient	Healthy adults 18 to 50 years of age
Subjects	329
	Arm A: pentamer (low)/gB(low)/adjuvant vaccine
	Arm B: pentamer (med)/gB(low)/adjuvant vaccine
Treatment arms	Arm C: pentamer (med)/gB(med)/adjuvant vaccine
	Arm D: pentamer (high)/gB(med)/adjuvant vaccine
	Arm F: placebo (saline)
Description	A randomised, observer-blind, placebo-controlled, dose escalation trial to assess safety, reactogenicity and immunogenicity of a candidate CMV vaccine comprising recombinant protein and adjuvant
T. I.	Trial start: Q4 2021
Timeline	Data anticipated: 2026+
Key end points	Safety, reactogenicity and immunogenicity
Clinicaltrials.gov	<u>Link</u>



GSK3943104 (Therapeutic HSV)

Phase	I/II	
Patient	Healthy participants aged 18-60 years negative for HSV-2	
	HSV-2 and HSV-1 patients with ≥3 episodes of GH in the previous	s year
Subjects	Part 1: 245; Part 2: 240	
	Arm A: non-adjuvanted HSV formulation 1 - part 1 group	Arm H: HSV formulation 2 with adjuvant 2 - part 1 group
	Arm B: non-adjuvanted HSV formulation 2 - part 1 group	Arm I: HSV formulation 3 with adjuvant 2 - part 1 group
	Arm C: non-adjuvanted HSV formulation 3 - part 1 group	Arm J: part 1 group (placebo)
Treatment arms	Arm D: HSV formulation 1 with adjuvant 1 - part 1 group	Arm K: selected formulation - part 2 group
	Arm E: HSV formulation 2 with adjuvant 1 - part 1 group	Arm L: selected formulation - part 2 group
	Arm F: HSV formulation 3 with adjuvant 1 - part 1 group	Arm M: part 2 group (placebo)
	Arm G: HSV formulation 1 with adjuvant 2 - part 1 group	
Description	An observer-blind, randomised, placebo-controlled, multi-country trial to evaluate reactogenicity, safety, immune response and efficacy of an HSV vaccine	
-	Trial start: Q1 2022	
Timeline	Data anticipated: 2026+	
	Part 1: Percentage of participants reporting each solicited admin	istration site event; dose selection
Key end points	Part 2: Clinical efficacy (TTFE)	
Clinicaltrials.gov	<u>Link</u>	



GSK4077164 (iNTS Typhimurium + Enteritidis)

Phase	I/IIa		
Patient	Healthy European and African adults		
Subjects	155		
	Arm A: iNTS-TCV low dose group - Europe	Arm F: Step 2 group (placebo) - Europe	
	Arm B: iNTS-GMMA and TCV low doses group - Europe	Arm G: iNTS-TCV full dose_2 group - Africa	
Treatment arms	Arm C: Step 1 group (placebo) - Europe	Arm H: iNTS-GMMA and TCV full doses_2 group - Africa	
	Arm D: iNTS-TCV full dose_1 group - Europe	Arm I: Stage 2 group (control) - Africa	
	Arm E: iNTS-GMMA and TCV full doses_1 group - Europe		
Description	An observer-blind, randomised, controlled, two-stage, multi-country trial to evaluate the safety, reactogenicity and immune response of the trivalent vaccine against iNTS and Typhoid fever		
Time alling	Trial start: Q3 2022		
Timeline	Data anticipated: H2 2024		
Key end points	To evaluate the safety, reactogenicity and immunogenicity profile of iNTS-TCV vaccine in healthy European/African adults		
Clinicaltrials. gov	<u>Link</u>		



GSK4077164 (iNTS S. typhimurium + S. enteritidis + S. Typhi)

Phase	Ila
Patient	Adults, children and infants, including dose-finding in infants in Africa (Ghana)
Subjects	20 adults/40 children/60 infants 9 months/ 396 infants 6 weeks
	Stage 1: Age-de-escalation
	Adults (dose C or control)
Treatment	Children (dose B or C or control)
arms	Infants, 9 months (dose A, B, C or control)
	Infants, 6 months (dose A, B, C, or control)
	Stage 2: Dose finding in infants 6 weeks of age
Description	An observer-blind, randomized, controlled, age-de-escalation, single center interventional study to evaluate the safety, reactogenicity, and immune response of the GVGH iNTS vaccine against <i>S. typhimurium</i> and <i>S. enteritidis</i> , in adults, children and infants, including dose-finding in infants, in Africa (Ghana)
T:	Trial start: Q1 2024
Timeline	Data anticipated: 2026+
Key end points	To evaluate the safety, reactogenicity and immunogenicity profile of iNTS-GMMA vaccine in adults, children and infants (Ghana)
Clinicaltrials. gov	<u>Link</u>



ganfeborole

Phase	IIb	
Patient	Adults with newly diagnosed, rifampicin-susceptible pulmonary TB	
Subjects	2500	
Treatment arms	Arm A: Isoniazid + rifampicin + pyrazinamide + ethambutol for 8 weeks then isoniazid + rifampicin for 16 weeks Arm B: Bedaquiline + delamanid + moxifloxacin Arm C: Bedaquiline + delamanid + moxifloxacin + ganfeborole Arm D: Bedaquiline + delamanid + pyrazinamide + ganfeborole Arm E: Bedaquiline + delamanid + linezolid (for first 8 weeks) + ganfeborole Arm F: Bedaquiline + pretomanid + moxifloxacin + ganfeborole	Arm G: Bedaquiline + delamanid + moxifloxacin + BTZ-043 Arm H: Bedaquiline + delamanid + pyrazinamide + BTZ-043 Arm I: Bedaquiline + delamanid + linezolid (for first 8 weeks) + BTZ-043 Arm J: Bedaquiline + pretomanid + moxifloxacin + BTZ-043 Arm K: Bedaquiline + moxifloxacin + pyrazinamide + BTZ-043 Arm L: Bedaquiline + delamanid + ganfeborole + BTZ-043
Description	A seamless phase 2B/C platform trial to evaluate multiple regimens and durations of treatment in pulmonary tuberculosis	
Timeline	Trial start: Q1 2024 Data anticipated: 2026+	
Key end points	Primary efficacy outcome in phase 2B, rate of change in log10 (time to positivity), primary efficacy outcome in phase 2C, the proportion of participants with a favourable outcome status	
Clinicaltrials.gov	<u>Link</u>	



GSK3536867 (Salmonella typhoid + paratyphoid A)

Phase	I
Patient	Healthy adults aged 18-50 years in Europe
Subjects	96
	Arm A: Step 1a low dose without adjuvant group
	Arm B: Step 1a control group
	Arm C: Step 1b low dose with adjuvant group
Treatment arms	Arm D: Step 1b control group
	Arm E: Step 2 full dose without adjuvant group
	Arm F: Step 2 full dose with adjuvant group
	Arm G: Step 2 control group
Description	An observer-blind, randomised, controlled, single-centre trial to evaluate the safety, reactogenicity and immune responses to an adjuvanted and non-adjuvanted conjugate vaccine against Salmonella Typhi and Salmonella Paratyphi A
T: I'	Trial start: Q4 2022
Timeline	Data anticipated: H2 2024
Key end points	Percentage of participants with solicited administration-site events, systemic events, unsolicited adverse event and any serious adverse events after the first vaccination
Clinicaltrials.gov	<u>Link</u>



GSK2556286 (Tuberculosis)

Phase	1
Patient	Healthy adults
Subjects	120
	Arm A: Part A - GSK2556286 with up to 11 cohorts
Treatment arms	Arm B: Part A - placebo
reatment arms	Arm C: Part B - GSK2556286 with up to 4 cohorts
	Arm D: Part B - placebo
Description	A randomised, double blind (sponsor unblinded), placebo-controlled, first time in human trial to evaluate the safety, tolerability and pharmacokinetics of single and repeat oral doses and the food effect of GSK2556286
Time aline	Trial start: Q4 2020
Timeline	Data anticipated: H2 2024
Key end points	SAEs and non-SAEs
Clinicaltrials.gov	<u>Link</u>



GSK4024484 (Malaria)

Phase	I and the second	
Patient	Healthy adults aged 18-60 years	
Subjects	144	
Treatment arms	Group/Arm 1: 6mg SAD GSK'484 or placebo (fasted state) Group/Arm 2: 12mg SAD GSK'484 or placebo (fasted state) Group/Arm 3: 24mg SAD GSK'484 or placebo (fasted state) Group/Arm 4: 40mg SAD GSK'484 or placebo (fasted state) Group/Arm 5: 60mg SAD GSK'484 or placebo (fasted state) Group/Arm 6: 80mg SAD GSK'484 or placebo (fasted state) Group/Arm 7: Food Effect (GSK'484 or placebo in fed state)	
Description	A randomised, double-blind placebo-controlled, First Time in Human Study to evaluate the safety and pharmacokinetics of single and multiple oral doses and food effect of GSK4024484	
Timeline	Trial start: Q4 2023 Data anticipated: 2025	
Key end points	Number of participants with AEs and SAEs	
Clinicaltrials.gov	<u>Link</u>	



GSK3882347 (Uncomplicated UTI)

Phase	Ib
Patient	Female participants with acute uncomplicated urinary tract infection
Subjects	80
	GSK3882347
Treatment arms	Nitrofurantoin
Description	A double-blind, double dummy, randomised, nitrofurantoin controlled, repeat oral dose trial to investigate the safety, tolerability, pharmacokinetics and microbiological response of GSK3882347 in female participants with acute uncomplicated urinary tract infection
Timeline	Trial start: Q4 2022
Imeline	Data anticipated: 2025
Key end points	Numbers of participants with microbiological response (responder/non-responder of GSK3882347) at the TOC visit
Clinicaltrials.gov	<u>Link</u>



GSK3923868 (Rhinovirus disease)

Phase	Ib
Patient	Participants with mild asthma
Subjects	48
T	Arm A: GSK3923868
Treatment arms	Arm B: placebo
Description	A randomised, double-blind, placebo controlled, repeat dose trial to assess the efficacy, safety, tolerability, pharmacokinetics and pharmacodynamics of inhaled GSK3923868 during experimental human rhinovirus infection participants with mild asthma
Time aline	Trial start: Q2 2022
Timeline	Trial end: Q2 2024
Key end points	AUC of CfB in LRTS score from day of inoculation up to discharge
Clinicaltrials.gov	<u>Link</u>



GSK3965193 (Chronic HBV infection)

Phase	1/11
Patient	Healthy participants and those living with chronic hepatitis B infection
Subjects	132
Treatment arms	Part 1 cohort 1: GSK3965193 and placebo Part 1 cohort 2: GSK3965193 and placebo Part 2A cohort 3: GSK3965193 or placebo Part 2A cohort 4: GSK3965193 or placebo Part 2A cohort 5: GSK3965193 or placebo Part 2B cohort 6: GSK3965193 Part 3 cohort 7: GSK3965193 or placebo Part 4 cohort 8: GSK3965193 and bepirovirsen or placebo and bepirovirsen
Description	Four-part, randomised, double-blind (Parts 1, 2A, 3 and 4), multi-centre, placebo-controlled trial to assess the safety, tolerability, pharmacokinetics and pharmacodynamics of GSK3965193 monotherapy in healthy participants and in participants living with chronic hepatitis B infection; and GSK3965193 in combination with bepirovirsen
Timeline	Trial start: Q2 2022 Data anticipated: 2026+
Key end points	Number of participants with AEs, SAEs, and withdrawals due to AEs Part 3: Change from Baseline in HBsAg levels Part 4: Number of participants achieving sustained virologic response
Clinicaltrials.gov	<u>Link</u>



HIV



HIV

VH3810109

NCT05996471 - EMBRACE

Phase IIb	
Patient An	tiretroviral therapy (ART)-experienced adults living with HIV
Subjects 124	4
Gro	oup 1: VH3810109 + cabotegravir
Treatment arms	oup 2 VH3810109 + rHuPH20 + cabotegravir
Gro	oup 3: Active comparator - Participants receiving standard of care (SOC) tiretroviral therapy (ART)
Description and into	multicentre, randomised, open-label, trial comparing the efficacy, safety, PK, d tolerability of VH3810109, administered either intravenously or as a boutaneous infusion with rHuPH20, in combination with cabotegravir given ramuscularly, to standard of care in virologically suppressed, antiretroviral erapy (ART)-experienced adults living with HIV
	al start: Q3 2023
Timeline Da	ata anticipated: H2 2024
Key end points Sat	fety, plasma HIV-1 levels
Clinicaltrials.gov <u>Lin</u>	n <u>k</u>





Phase	Ila
Patient	Treatment-naïve adults living with HIV-1
Subjects	28
Treatment arms	Arm A: VH3738837 Arm B: placebo
Description	A randomized, double-blind (sponsor-unblinded), placebo-controlled, adaptive study to investigate the antiviral effect, safety, tolerability and pharmacokinetics of VH3739937 in treatment-naïve adults living with HIV-1
Timeline	Trial start: Q1 2024 Data anticipated: H2 2024
Key end points	AEs and SAEs, concentrations of VH3738837
Clinicaltrials.gov	<u>Link</u>





VH4004280 & VH4011499

NCT06012136

Phase	1
Patient	Healthy adults
Subjects	160
Treatment arms	Arm A: VH4004280 Arm B: Placebo
	Arm C: VH4011499
Description	A double-blind (sponsor-unblinded), placebo-controlled, randomized, single dose escalation study to evaluate the safety, tolerability, and pharmacokinetics of a parenterally administered suspension of investigational capsid inhibitors in healthy adults
Timeline	Trial start: Q3 2023
	Data anticipated: 2025+
Key end points	AEs, PK
Clinicaltrials. gov	<u>Link</u>

NCT06039579 - CINNAMON

Phase	II
Patient	HIV-1 infected treatment-naïve adults
Subjects	42
	Arm A: VH4004280
Treatment	Arm B: VH4011499
arms	Arm C: VH4004280-matching placebo
	Arm D: VH4011499-matching placebo
Description	A randomized, double-blind (sponsor-unblinded), placebo-controlled trial to investigate the antiviral effect, safety, tolerability and pharmacokinetics of orally administered investigational capsid inhibitor monotherapy in HIV-1 infected treatment-naïve adults
Timeline	Trial start anticipated: H2 2023
Ilmeline	Data anticipated: H2 2024
Key end points	Maximum change from baseline (Day 1) in plasma HIV-1 RNA
Clinicaltrials. gov	<u>Link</u>





VH4524184

Phase	Ila
Patient	HIV-1 infected treatment naïve adults
Subjects	28
Treatment arms	Arm A: VH4524184 Arm B: Placebo
Description	A randomized, double-blind (sponsor unblinded), placebo-controlled study to investigate the antiviral effect, safety, tolerability and pharmacokinetics of VH4524184 in HIV-1 infected treatment naïve adults
Timeline	Trial start: Q1 2024 Data anticipated: H2 2024
Key end points	Maximum change from baseline in log10 plasma HIV-1 RNA
Clinicaltrials.gov	<u>Link</u>



HIV

cabotegravir

NCT05418868 NCT06033547

Phase	I	Phase	T. Company of the com
Patient	Healthy adult volunteers	Patient	Healthy adult volunteers
Subjects	60	Subjects	48
Treatment arms	Part A: Participants receiving CAB 200 mg/mL with rHuPH20 Part C: Participants receiving CAB 400 mg/mL	Treatment arms	Part A: Participants receiving cabotegravir Formulation F Part B: Participants receiving cabotegravir Formulation G
Description	Part D: Participants receiving CAB 400 mg/mL with rHuPH20 A multi-centre, open-label, single dose escalation trial to evaluate the pharmacokinetics, safety and tolerability of long-acting cabotegravir co-administered with recombinant human hyaluronidase PH20 (rHuPH20) in healthy adult volunteers	 Description	An open-label, single dose escalation study to evaluate the pharmacokinetics, safety and tolerability of two different formulations of long-acting cabotegravir administered to healthy adult participants
		Timeline	Trial start: Q3 2023
-	Trial start: Q2 2022		Data anticipated: 2025
Timeline	Data anticipated: 2025	Key end points	Plasma concentrations of cabotegravir
Key end points	Plasma concentrations of cabotegravir	Clinicaltrials.gov	<u>Link</u>
Clinicaltrials.gov	<u>Link</u>		





Nucala (mepolizumab)

NCT04133909 - MATINEE

Phase	III	
Patient	Participants with chronic obstructive pulmonary disease (COPD) experiencing frequent exacerbations and characterised by eosinophil levels	
Subjects	806	
Treatment arms	Arm A: placebo	
	Arm B: mepolizumab	
Description	A multicentre randomised, double-blind, parallel-group, placebo-controlled trial of mepolizumab 100 mg subcutaneously as add-on treatment in participants with COPD experiencing frequent exacerbations and characterised by eosinophil levels	
Timeline	Trial start: Q4 2019	
Imeline	Data anticipated: H2 2024	
Key end points	Annualised rate of moderate or severe exacerbations	
Clinicaltrials.gov	<u>Link</u>	



NCT04719832 - SWIFT-1

NCT04718103 - SWIFT-2

Phase	III	Phase	III
Patient	Adult and adolescents with severe uncontrolled asthma with an eosinophilic phenotype	Patient	Adult and adolescents with severe uncontrolled asthma with an eosinophilic phenotype
Subjects	395	Subjects	397
Treatment arms	Arm A: depemokimab plus SoC	Treatment arms	Arm A: depemokimab plus SoC
	Arm B: placebo plus SoC	reatment arms	Arm B: placebo plus SoC
Description	A 52-week, randomised, double-blind, placebo-controlled, parallel-group, multi-centre trial of the efficacy and safety of depemokimab adjunctive therapy in adult and adolescent participants with severe uncontrolled asthma with an eosinophilic phenotype	Description	A 52-week, randomised, double-blind, placebo-controlled, parallel-group, multi-centre trial of the efficacy and safety of depemokimab adjunctive therapy in adult and adolescent participants with severe uncontrolled asthma with an eosinophilic phenotype
T: I:	Trial start: Q1 2021	Timeline	Trial start: Q1 2021
Timeline	Data reported: Q2 2024		Data reported: Q2 2024
Key end points	Annualised rate of clinically significant exacerbations over 52 weeks	Key end points	Annualised rate of clinically significant exacerbations over 52 weeks
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>
		-	



NCT05243680 - AGILE

Phase	III
Patient	Adult and adolescents with severe asthma with an eosinophilic phenotype from studies SWIFT-1 and SWIFT-2
Subjects	637
Treatment arms	Participants diagnosed with asthma receiving depemokimab
Description	A 52-week, open label extension phase of SWIFT-1 and SWIFT-2 to assess the long-term safety and efficacy of depemokimab adjunctive therapy in adult and adolescent participants with severe uncontrolled asthma with an eosinophilic phenotype
Timeline	Trial start: Q1 2022
imeiine	Data anticipated: 2025
Key end points	Number of participants with AEs and SAEs and incidence of immunogenicity over 52 weeks
Clinicaltrials.gov	<u>Link</u>

NCT04718389 - NIMBLE

Phase	III
Patient	Adult and adolescent severe asthmatic participants with an eosinophilic phenotype treated with depemokimab compared with mepolizumab or benralizumab
Subjects	1700
T	Arm A: participants receiving depemokimab plus placebo matching prior anti- IL-5/5R treatment
Treatment arms	Arm B: participants receiving prior anti-IL-5/5R treatment plus placebo matching depemokimab
Description	A 52-week, randomised, double-blind, double-dummy, parallel group, multicentre, non-inferiority trial assessing exacerbation rate, additional measures of asthma control and safety in adult and adolescent severe asthmatic participants with an eosinophilic phenotype treated with depemokimab compared with mepolizumab or benralizumab
T'	Trial start: Q1 2021
Timeline	Data anticipated: 2025
Key end points	Annualised rate of clinically significant exacerbations over 52 weeks
Clinicaltrials.gov	<u>Link</u>



NCT05274750 - ANCHOR-1

NCT05281523 - ANCHOR-2

Phase	III	Phase	III
Patient	Adults with chronic rhinosinusitis with nasal polyps (CRSwNP)	Patient	Adults with chronic rhinosinusitis with nasal polyps (CRSwNP)
Subjects	276	Subjects	264
Treatment arms	Arm A: depemokimab	Treatment arms	Arm A: depemokimab
reatment arms	Arm B: placebo		Arm B: placebo
Description	A randomized, double-blind, parallel group trial to assess the efficacy and safety of 100 mg subcutaneous depemokimab in patients with CRSwNP	Description	A randomized, double-blind, parallel group trial to assess the efficacy and safety of 100 mg subcutaneous depemokimab in patients with CRSwNP
Timeline	Trial start: Q2 2022	Timeline	Trial start: Q2 2022
rimeline	Data anticipated: H2 2024	i inieline	Data anticipated: H2 2024
	Change from baseline in total endoscopic nasal polyps (NP) score at week 52		Change from baseline in total endoscopic nasal polyps (NP) score at week 52
Key end points	Change from baseline in mean nasal obstruction verbal response scale (VRS) score from Week 49 through to Week 52	Key end points	Change from baseline in mean nasal obstruction verbal response scale (VRS) score from Week 49 through to Week 52
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>



NCT05263934 - OCEAN

Phase	III
Patient	Adults with relapsing or refractory eosinophilic granulomatosis with polyangiitis (EGPA) receiving standard of care therapy
Subjects	160
Treatment arms	Arm A: depemokimab + placebo matching mepolizumab
	Arm B: mepolizumab + placebo matching depemokimab
Description	A 52-week randomised, double-blind, double-dummy, parallel-group, multicentre, non-inferiority trial to investigate the efficacy and safety of depemokimab compared with mepolizumab in adults with relapsing or refractory EGPA receiving standard of care therapy
Timeline	Trial start: Q3 2022
	Data anticipated: 2025
Key end points	Number of participants with remission
Clinicaltrials.gov	<u>Link</u>

NCT05334368 - DESTINY

III
Adults with hypereosinophilic syndrome (HES) receiving standard of care therapy
120
Arm A: depemokimab
Arm B: placebo
A randomised, double-blind, placebo-controlled trial to investigate the efficacy and safety of depemokimab in adults with HES
Trial start: Q3 2022
Data anticipated: 2026+
Frequency of HES flares
<u>Link</u>



Respiratory/Immunology camlipixant

NCT05599191 - CALM-1

NCT05600777 - CALM-2

Phase	III	Phase	III
Patient	Adult participants with refractory chronic cough, including unexplained chronic cough	Patient	Adult participants with refractory chronic cough, including unexplained chronic cough
Subjects	825	Subjects	825
	Arm A: camlipixant 25 mg twice a day		Arm A: camlipixant 25 mg twice a day
Treatment arms	Arm B: camlipixant 50 mg twice a day	Treatment arms	Arm B: camlipixant 50 mg twice a day
	Placebo twice a day		Placebo twice a day
Description	A 52-week, randomised, double-blind, placebo-controlled, parallel-arm efficacy and safety study with open-label extension of camlipixant in adult participants with refractory chronic cough, including unexplained chronic cough	Description	A 24-week, randomised, double-blind, placebo-controlled, parallel-arm efficacy and safety study with open-label extension of camlipixant in adult participants with refractory chronic cough, including unexplained chronic cough
T. I.	Trial start: Q4 2022	Time aline	Trial start: Q1 2023
Timeline	Data anticipated: 2025		Data anticipated: 2025
Key end points	24-hour cough frequency	Key end points	24-hour cough frequency
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>



Respiratory/Immunology Ventolin (low carbon version of MDI)

Patient Participants aged 12 years and above with asthma Subjects 412 Treatment arms Arm A: Salbutamol HFA-134a Arm B: Salbutamol HFA-152a A randomized, double-blind, parallel group, multi-center study to evaluate the long-term safety of salbutamol rescue medication when administered via
Treatment arms Arm A: Salbutamol HFA-134a Arm B: Salbutamol HFA-152a A randomized, double-blind, parallel group, multi-center study to evaluate the long-term safety of salbutamol rescue medication when administered via
Arm B: Salbutamol HFA-152a A randomized, double-blind, parallel group, multi-center study to evaluate the long-term safety of salbutamol rescue medication when administered via
Arm B. Salbutamor AFA-132a A randomized, double-blind, parallel group, multi-center study to evaluate the long-term safety of salbutamol rescue medication when administered via
Description long-term safety of salbutamol rescue medication when administered via
metered dose inhalers containing the propellant HFA-152a or reference HFA- 134a
Trial start: Q2 2024 Timeline
Data anticipated: 2025
Key end points AEs
Clinicaltrials. gov



Respiratory/Immunology Benlysta (belimumab)

Phase	11/111
Patient	Adults with systemic sclerosis associated interstitial lung disease (SSc-ILD)
Subjects	300
Treatment arms	Arm A: belimumab + standard therapy
	Arm B: placebo + standard therapy
Description	A randomized, double-blind, placebo-controlled, parallel-group trial to evaluate the efficacy and safety of belimumab administered subcutaneously in adults with SSc-ILD
Timeline	Trial start: Q4 2023
	Data anticipated: 2026+
Key end points	Absolute change from baseline in Forced Vital Capacity (FVC) millilitre (mL) at week 52
Clinicaltrials.gov	<u>Link</u>



GSK3858279 (Osteoarthritis pain)

NCT05838742 - MARS-17

NCI	05838	/55 -	MEDI	UNE-I/

Phase	II
Patient	Adult participants with moderate to severe pain due to knee osteoarthritis
Subjects	420
Treatment arms	Arm A: GSK3858279 dose 1
	Arm B: GSK3858279 dose 2
	Arm C: GSK3858279 dose 3
	Arm D: GSK3858279 dose 4
	Arm E: placebo
Description	A multicentre, randomised, double-blind, placebo controlled, dose-finding trial of GSK3858279 in adult participants with moderate to severe pain due to knee osteoarthritis
Timeline	Trial start: Q4 2023
	Data anticipated: 2025
Key end points	Change from baseline in the weekly average of average daily knee pain intensity at week 12, assessed on Numeric Rating Scale (NRS)
Clinicaltrials.gov	<u>Link</u>

Phase	II
Patient	Adult participants with chronic diabetic peripheral neuropathic pain (DPNP)
Subjects	240
Treatment arms	Arm A: GSK3858279 dose 1
	Arm B: GSK3858279 dose 2
	Arm C: placebo
Description	A multicentre, randomised, double-blind, placebo-controlled trial to evaluate efficacy, safety, tolerability, pharmacokinetics and target engagement of GSK3858279 in adult participants with chronic DPNP
Timeline	Trial start: Q4 2023
	Data anticipated: 2025
Key end points	Change from baseline in the weekly average of average daily pain intensity at week 12, assessed on Numeric Rating Scale (NRS)
Clinicaltrials.gov	<u>Link</u>



GSK1070806 (Atopic dermatitis)

Phase	Пр
Patient	Patients with moderate to severe atopic dermatitis
Subjects	175
Treatment arms	Arm A: GSK1070806 dose 1
	Arm B: GSK1070806 dose 2
	Arm C: GSK1070806 dose 3
	Arm D: GSK1070806 dose 4
	placebo
Description	A randomized, double-blind, parallel group, placebo-controlled dose finding study to evaluate the efficacy, safety, pharmacokinetics, and pharmacodynamics of GSK1070806 SC injection
Timeline	Trial start: Q4 2023
	Data anticipated: 2025
Key end points	Percent change from baseline in eczema area and severity index (EASI) at Week 16
Clinicaltrials.gov	<u>Link</u>



GSK4527226 (Alzheimer's disease)

NCT06079190 - PROGRESS-AD

Phase	II
Patient	Participant must be in the Alzheimer's continuum as defined by the 2018 National Institute on Aging and Alzheimer's Association (NIAAA) Research Framework corresponding to the clinical categories of MCI due to AD and mild AD dementia.
Subjects	282
	Arm 1: GSK4527226 Dose 1
Treatment arms	Arm 2 GSK4527226 Dose 2
	Arm 3: Placebo
Description	A parallel group, randomized, double-blind, placebo-controlled, 3-arm, multicenter treatment study to evaluate the efficacy and safety of GSK4527226 [AL101] intravenous infusion compared with placebo in patients with early Alzheimer's Disease
Timeline	Trial start: Q4 2023
	Primary data reported: 2026+
Key end points	CDR-SB, iADRS, ADAS-Cog14, ADCS-ADL-MCI, ADCS-iADL, ADCOMS
Clinicaltrials.gov	<u>Link</u>



GSK3915393 (Pulmonary fibrosis)

Phase	II .
Patient	Participants with Idiopathic Pulmonary Fibrosis (IPF)
Subjects	150
Treatment arms	Arm A: GSK3915393
	Arm B: placebo
Description	A randomized, double-blind, placebo controlled, parallel group study (TRANSFORM) to evaluate the efficacy and safety of GSK3915393 in participants With Idiopathic Pulmonary Fibrosis (IPF)
Timeline	Trial start anticipated: Q2 2024
	Data anticipated: 2026+
Key end points	Absolute change from baseline in Forced Vital Capacity (FVC) in milliliters (mL) at Week 26
Clinicaltrials.gov	<u>Link</u>



Respiratory/Immunology GSK3862995 (COPD)

Phase	1
Patient	Part A: Healthy participants Part B: Participants with Chronic Obstructive Pulmonary Disorder
Subjects	130
Treatment arms	Part A: Single ascending dose (SAD) of GSK3862995B
	Part B, arm A: Repeat doses GSK3862995B
	Part B, arm B: Placebo
Description	A two-part randomized, double-blind, placebo-controlled study to investigate safety, tolerability, immunogenicity, pharmacokinetics and pharmacodynamics of GSK3862995B following single ascending doses in healthy participants and repeat doses in participants with Chronic Obstructive Pulmonary Disease (COPD)
Timeline	Trial start: Q4 2023
	Data anticipated: 2026+
Key end points	AEs and SAEs
Clinicaltrials.gov	<u>Link</u>



GSK4347859 (Systemic lupus erythematosus)

Phase	I
Patient	Healthy participants
Subjects	44
Treatment arms	Part 1, cohort 1: GSK4347859 or placebo
	Part 1, cohort 2: GSK4347859 or placebo
	Part 2, cohort 3: GSK4347859 (dose level A) or placebo
	Part 2, cohort 4: GSK4347859 (dose level B) or placebo
	Part 2, cohort 5: GSK4347859 (dose level C) or placebo
Description	A randomized, double-blind, placebo-controlled study to evaluate the safety, tolerability, pharmacokinetics and pharmacodynamics of GSK3996401 following single and multiple ascending doses of GSK4347859 in healthy participants
Timeline	Trial start: Q1 2024
	Data anticipated: 2025
Key end points	AEs and SAEs Maximum observed plasma concentration (Cmax) of GSK3996401 following administration of GSK4347859
Clinicaltrials.gov	<u>Link</u>



Respiratory/Immunology belantamab

Phase	Ib
Patient	Participants with moderate to severe Systemic Lupus Erythematosus (SLE)
Subjects	16
Treatment arms	belantamab
Description	A dose escalation, open label study to evaluate the safety, tolerability, pharmacokinetics and pharmacological effect of a single intravenous infusion of belantamab in participants with moderate to severe SLE
Timeline	Trial start anticipated: H2 2024
	Data anticipated: 2025
Key end points	AEs, SAEs
Clinicaltrials.gov	<u>Link</u>





Oncology Jemperli (dostarlimab)

NCT03981796 - RUBY ENGOT-EN6 GOG-3031

Phase	III
Patient	Patients with recurrent or primary advanced endometrial cancer
Subjects	785
	Arm A: dostarlimab + SoC followed by dostarlimab
Treatment arms	Arm B: placebo + SoC followed by placebo
i reatment arms	Arm C: dostarlimab + SoC followed by dostarlimab+niraparib
	Arm D: placebo (+SoC) followed by placebo
Description	A randomised, double-blind, multi-centre trial of dostarlimab plus carboplatin- paclitaxel with and without niraparib maintenance versus placebo plus carboplatin-paclitaxel in patients with recurrent or primary advanced endometrial cancer
Timeline	Trial start: Q3 2019
	Part 1 data reported: Q4 2022; Part 2 data reported: Q4 2023
Key end points	Part 1: Co-primary PFS by IA (dMMR/MSI-H and ITT) and OS (ITT)
	Part 2: Primary PFS (ITT) and key secondary OS (ITT)
Clinicaltrials.gov	<u>Link</u>

NCT04581824 - PERLA

Phase	II .
Patient	Participants with metastatic non-squamous non-small cell lung cancer (NSCLC)
Subjects	244
Treatment arms	Arm A: dostarlimab + chemotherapy Arm B: pembrolizumab + chemotherapy
Description	A randomised, double-blind trial to evaluate the efficacy of dostarlimab plus chemotherapy versus pembrolizumab plus chemotherapy in metastatic non-squamous NSCLC
Timeline	Trial start: Q4 2020 Primary data reported: Q4 2022
Key end points	ORR, OS, PFS
Clinicaltrials.gov	<u>Link</u>



Jemperli (dostarlimab)

NCT02715284 - GARNET

Phase	I/II
Patient	Participants with advanced solid tumors
Subjects	740
Treatment arms	Part 1: dostarlimab at ascending weight doses Part 2A: dostarlimab fixed dose of 500mg Q3W or 1000mg administered Q6W dose Part 2B: Cohort A1 dMMR/MSI-H endometrial Part 2B: Cohort A2 MMR proficient/MSS endometrial Part 2B: Cohort E: NSCLC Part 2B: Cohort F non-endometrial dMMR/MSI-H & POLE-mutation Part 2B: Cohort G PROC without known BRCA
Description	A multi-centre, open-label, first-in-human trial evaluating dostarlimab in participants with advanced solid tumors who have limited available treatment options
Timeline	Trial start: Q1 2016 Primary data reported: Q1 2019
Key end points	ORR, DoR, safety
Clinicaltrials.gov	<u>Link</u>

NCT05723562 - AZUR-1

Phase	II
Patient	Patients with untreated stage II/III mismatch repair deficient/high microsatellite instability (dMMR/MSI-H) locally advanced rectal cancer
Subjects	150
Treatment arms	dostarlimab monotherapy
Description	A single-arm, open-label trial with dostarlimab monotherapy in participants with untreated stage II/III dMMR/MSI-H locally advanced rectal cancer
Timeline	Trial start: Q1 2023
	Data anticipated: 2026+
Key end points	Sustained cCR for 12, 24 and 36 months, EFS at 3 years
Clinicaltrials.gov	<u>Link</u>



Oncology Jemperli (dostarlimab)

NCT05855200 - AZUR-2

Phase	III
Patient	Participants with untreated T4N0 or Stage III (resectable), mismatch repair deficient/high microsatellite instability (dMMR/MSI-H) colon cancer
Subjects	711
	Arm A: dostarlimab
Treatment arms	Arm B: Standard of care (FOLFOX/CAPEOX) or expectant observation post surgery.
Description	An open-label, randomized trial of perioperative dostarlimab monotherapy versus standard of care in participants with untreated T4N0 or Stage III dMMR/MSI-H resectable colon cancer
Timeline	Trial start: Q3 2023
	Data anticipated: 2026+
Key end points	EFS assessed by Blinded Independent Central Review (BICR)
Clinicaltrials.gov	<u>Link</u>

NCT06256588 - JADE

Phase	III
Patient	Participants have newly diagnosed unresected locally advanced histologically confirmed HNSCC of the oral cavity, oropharynx, hypopharynx or larynx and completed cisplatin plus radiotherapy (termed "CRT" in this protocol) with curative intent and has no evidence of distant metastatic disease.
Subjects	864
Treatment arms	Arm A: dostarlimab Arm B: Placebo
Description	A randomized, double-blind, placebo-controlled study to evaluate dostarlimab as sequential therapy after chemoradiation in participants with locally advanced unresected head and neck squamous cell carcinoma
Timeline	Trial start: Q1 2024
	Data anticipated: 2026+
Key end points	EFS assessed by Blinded Independent Central Review (BICR)
Clinicaltrials.gov	<u>Link</u>



Oncology Zejula (niraparib)

NCT03602859 - FIRST

Phase	III
Patient	Participants with Stage III or IV nonmucinous epithelial ovarian cancer
Subjects	1402
	Arm A: SOC (carboplatin + paclitaxel ± bevacizumab) +placebo
Treatment arms	Arm B: SOC + niraparib
	Arm C: SOC + dostarlimab + niraparib
Description	A randomised, double-blind comparison of platinum-based therapy with TSR-042 and niraparib versus standard of care platinum-based therapy as first-line treatment of Stage III or IV nonmucinous epithelial ovarian cancer
Time a 15m a	Study start: Q4 2018
Timeline	Data anticipated: H2 2024
Key end points	PFS for PD-L1 positive participants. Primary analysis is ARM B vs ARM C.

NCT04475939 - ZEAL-1L

Phase	III
Patient	Participants whose disease has remained stable or responded to 1L platinum-based chemo with pembrolizumab for stage IIIB/IIIC or IV NSCLC
Subjects	666
Tue ortine and ordine	Arm A: niraparib plus pembrolizumab
Treatment arms	Arm B: placebo plus pembrolizumab
Description	A randomised, double-blind, placebo-controlled, multicentre study comparing niraparib plus pembrolizumab versus placebo plus pembrolizumab as maintenance therapy
Timeline	Study start: Q4 2020
	Data anticipated: H2 2024
Key end points	OS, PFS assessed by BICR using Response Evaluation Criteria in Solid Tumors (RECIST)
Clinicaltrials.gov	<u>Link</u>



Blenrep (belantamab mafodotin)

NCT04126200 - DREAMM-5

Phase	1/11
Patient	Participants with relapsed/refractory multiple myeloma (RRMM)
Subjects	464
	Substudy 1: belantamab mafodotin + OX40 (GSK3174998)
	Substudy 2: belanatamab mafodotin + feladilimab
	Substudy 3: belantamab mafodotin + nirogacestat (GSI)
Treatment arms	Substudy 4: belantamab mafodotin + dostarlimab
	Substudy 5: belantamab mafodotin + isatuximab
	Substudy 6: belantamab mafodotin + nirogacestat + lenalidomide + dexamethasone
	Substudy 7: belantamab mafodotin + nirogacestat + pomalidomide + dexamethasone
Description	A randomised, open-label platform trial utilizing a master protocol to trial belantamab mafodotin as monotherapy and in combination with anti-cancer treatments
· ··	Trial start: Q4 2019
Timeline	Data anticipated: 2026+
Key end points	Dose escalation phase: DLT, safety, ORR Cohort expansion phase: ORR, CBR, safety
Clinicaltrials.gov	<u>Link</u>

NCT03544281 - DREAMM-6

Phase	1/11
Patient	Participants with relapsed/refractory multiple myeloma (RRMM)
Subjects	152
Tue salar and sugar	Arm A: belantamab mafodotin + lenalidomide + dexamethasone
Treatment arms	Arm B: belantamab mafodotin + bortezomib + dexamethasone
Description	An open-label, dose escalation and expansion trial to evaluate safety, tolerability and clinical activity of the antibody-drug conjugate belantamab mafodotin administered in combination with lenalidomide plus dexamethasone (Arm A), or bortezomib plus dexamethasone (Arm B)
Time alim a	Trial start: Q3 2018
Timeline	Trial end: Q1 2024
Key end points	DLT, safety, ORR, PK
Clinicaltrials.gov	<u>Link</u>



Blenrep (belantamab mafodotin)

NCT04246047 - DREAMM-7

Phase	III
Patient	Participants with relapsed/refractory multiple myeloma (RRMM)
Subjects	571
Treatment arms	Arm A: belantamab mafodotin + bortezomib + dexamethasone (B-Vd)
	Arm B: daratumumab, bortezomib + dexamethasone (D-Vd)
Description	A multicentre, open-label, randomised trial to evaluate the efficacy and safety of the combination of belantamab mafodotin, bortezomib and dexamethasone (B-Vd) compared with the combination of daratumumab, bortezomib and dexamethasone (D-Vd)
Timeline	Trial start: Q2 2020
	Primary data reported: Q4 2023
Key end points	PFS, CRR, ORR, DoR, TTR, TTP, OS, PFS2, MRD negativity rate, safety
Clinicaltrials.gov	<u>Link</u>

NCT04246047 - DREAMM-8

Phase	III
Patient	Participants with relapsed/refractory multiple myeloma (RRMM)
Subjects	300
Tue situe and signed	Arm A: belantamab mafodotin+ pomalidomide + dexamethasone (B-Pd)
Treatment arms	Arm B: Pomalidomide, bortezomib + dexamethasone (P-Vd)
Description	A multicentre, open-label, randomised trial to evaluate the efficacy and safety of belantamab mafodotin in combination with pomalidomide and dexamethasone (B-Pd) versus pomalidomide plus bortezomib and dexamethasone (PVd)
Timeline	Trial start: Q4 2020
Timeline	Primary data reported: Q1 2024
Key end points	PFS, MRD negativity rate, ORR, CRR, VGPR or better rate, DoR, TTBR, TTR, TTP, OS, PFS2, safety
Clinicaltrials.gov	<u>Link</u>



Blenrep (belantamab mafodotin)

NCT04091126 - DREAMM-9

Phase	1
Patient	Patients with newly diagnosed multiple myeloma (MM)
Subjects	144
	Belantamab mafodotin, selected doses
	Bortezomib, administered subcutaneously or intravenously approximately 1 hour after the belantamab mafodotin infusion until Cycle 8
Treatment arms	Lenalidomide, administered as 25 or 10 mg orally, depending upon renal function.
	Dexamethasone, administered orally as 20 mg in cycles 1-8 and 40 mg in Cycle 9 onwards
Description	A randomised, dose and schedule evaluation trial to investigate the safety, pharmacokinetics, pharmacodynamics and clinical activity of belantamab mafodotin administered in combination with standard of care
Timeline	Trial start: Q4 2019
	Data anticipated: 2025
Key end points	DLT, safety, RDI of lenalidomide and bortezomib, PK, PD, ORR, CRR, VGPR or better
Clinicaltrials.gov	<u>Link</u>

NCT04398745 - DREAMM-12

Phase	
Patient	Relapsed/refractory multiple myeloma (RRMM) who have normal and varying degrees of impaired renal function
Subjects	36
Treatment arms	belantamab mafodotin monotherapy
Description	A trial to evaluate the pharmacokinetics and safety of belantamab mafodotin monotherapy
Timeline	Trial start: Q4 2020
	Data anticipated: 2025
Key end points	PK, change in vital signs, safety
Clinicaltrials.gov	<u>Link</u>



Blenrep (belantamab mafodotin)

NCT04398680 - DREAMM-13

Phase	I e
Patient	Relapsed/refractory multiple myeloma (RRMM) who have normal and impaired hepatic function
Subjects	28
Treatment arms	belantamab mafodotin monotherapy
Description	A trial to evaluate the pharmacokinetics and safety of belantamab mafodotin monotherapy in participants who have normal and impaired hepatic function
Timeline	Trial start: Q2 2021
	Data anticipated: 2025
Key end points	PK, change in vital signs, safety
Clinicaltrials.gov	<u>Link</u>
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NCT05064358 - DREAMM-14

Phase	
Patient	Participants with relapsed/refractory multiple myeloma (RRMM)
Subjects	180
Treatment arms	belantamab mafodotin
Description	A randomised, parallel, open-label study to investigate the safety, efficacy and pharmacokinetics of various dosing regimens of single-agent belantamab mafodotin (GSK2857916)
Timeline	Study start: Q1 2022 Data anticipated: H2 2024
Key end points	% of patients with >= Gr 2 ocular events, safety, ORR, TTR, DoR, TTP, PFS, OS
Clinicaltrials.gov	<u>Link</u>



Oncology cobolimab

NCT04655976 - COSTAR LUNG

Phase	11/111
Patient	Patients with advanced non-small cell lung cancer (NSCLC) who have progressed on prior anti-PD-(L)1 therapy and chemotherapy
Subjects	750
	Arm A: cobolimab + dostarlimab + docetaxel
Treatment arms	Arm B: dostarlimab + docetaxel
	Arm C: docetaxel
Description	A randomised, open label trial comparing cobolimab + dostarlimab + docetaxel to dostarlimab + docetaxel to docetaxel alone
Timeline	Trial start: Q4 2020
	Data anticipated: 2025
Key end points	OS, ORR, PFS, DoR, TTD
Clinicaltrials.gov	<u>Link</u>



belrestotug & CD226 assets

NCT05565378 - GALAXIES LUNG-201

Phase	II
Patient	Participants with previously untreated, locally advanced/metastatic, Programmed Death Ligand 1-selected non small cell lung cancer (NSCLC)
Subjects	300
Treatment arms	Comparator Arm: pembrolizumab monotherapy Intervention Arm: dostarlimab monotherapy Substudy 1A: dostarlimab + belrestotug (Dose A) Substudy 1B: dostarlimab + belrestotug (Dose B) Substudy 1C: dostarlimab + belrestotug (Dose C) Substudy 2: dostarlimab + belrestotug + nelistotug
Description	A randomized, open-label, platform trial utilizing a master protocol to evaluate novel immunotherapy combinations in participants with previously untreated, locally advanced/metastatic, Programmed Death Ligand 1-selected NSCLC
Timeline	Trial start: Q4 2022 Data anticipated: 2025
Key end points	ORR
Clinicaltrials.gov	<u>Link</u>

NCT06472076 - GALAXIES LUNG-301

Phase	III
Patient	Participants with previously untreated, unresectable, locally advanced or metastatic PD-L1 selected non-small cell lung cancer
Subjects	1000
Treatment	Experimental: dostarlimab plus belrestotug
arms	Comparator: pembrolizumab plus placebo
Description	A randomized, multicenter, double-blind trial to investigate the safety and efficacy of belrestotug in combination with dostarlimab compared with placebo in combination with pembrolizumab in participants with previously untreated, unresectable, locally advanced or metastatic PD-L1 selected non-small cell lung cancer
Timeline	Trial start: Q3 2024
	Data anticipated: 2026+
Key end points	PFS, OS
Clinicaltrials. gov	<u>Link</u>



belrestotug & CD226 assets

NCT03739710 - ENTRÉE Lung

Patient Participants with non-small cell lung cancer (NSCLC) Subjects 185 Treatment arms Arm B: dostarlimab + belrestotug Arm C: dostarlimab + belrestotug + nelistotug Description A randomized, open-label platform trial utilizing a master protocol to trial novel regimens versus standard of care treatment in NSCLC participants Trial start: Q1 2019 Data anticipated: 2025 Part 1: Number of participants with AEs, SAEs, DLT, clinically significant
Treatment arms Arm B: dostarlimab + belrestotug Arm C: dostarlimab + belrestotug + nelistotug Description A randomized, open-label platform trial utilizing a master protocol to trial novel regimens versus standard of care treatment in NSCLC participants Trial start: Q1 2019 Data anticipated: 2025
Treatment arms Arm C: dostarlimab + belrestotug + nelistotug Description A randomized, open-label platform trial utilizing a master protocol to trial novel regimens versus standard of care treatment in NSCLC participants Trial start: Q1 2019 Data anticipated: 2025
novel regimens versus standard of care treatment in NSCLC participants Trial start: Q1 2019 Data anticipated: 2025
Timeline Data anticipated: 2025
<u> </u>
changes in vital signs, physical examination and laboratory parameters. Number of participants requiring dose modifications. Part 2: Overall survival
Clinicaltrials.gov Link

NCT06062420 - GALAXIES H&N-202

Phase	II
Patient	Participants with recurrent/metastatic PD-L1 positive squamous cell carcinoma of the head and neck
Subjects	360
	Arm A: dostarlimab monotherapy
Treatment arms	Arm B: dostarlimab and belrestotug
rreatment arms	Arm C: dostarlimab and nelistotug
	Arm D: dostarlimab and belrestotug and nelistotug
Description	A randomized, open-label, platform study using a master protocol to evaluate novel immunotherapy combinations as first-line treatment in participants with recurrent/metastatic PD-L1 positive squamous cell carcinoma of the head and neck
Timeline	Trial start: Q4 2023
	Data anticipated: 2026+
Key end points	ORR
Clinicaltrials.gov	<u>Link</u>



belrestotug & CD226 assets

NCT04446351 - nelistotug FTIH

Phase	1
Patient	Participants with advanced solid tumours
Subjects	244
	Arm A: nelistotug
	Arm B: nelistotug + dostarlimab
Treatment arms	Arm C: dostarlimab
rreatment arms	Arm D: dostarlimab + belrestotug
	Arm E: dostarlimab + belrestotug + nelistotug
	Arm G: dostarlimab + cobolimab
Description	A first time in human, open-label trial of nelistotug (GSK6097608) administered as monotherapy and in combination with anticancer agents
Timeline	Trial start: Q1 2020
	Data anticipated: 2025
Key end points	DLT, AEs and SAEs
Clinicaltrials.gov	<u>Link</u>

NCT05277051 - PVRIG FTIH

Phase

	<u> </u>
Patient	Participants with selected advanced solid tumors
Subjects	162
	Arm A: GSK4381562 monotherapy
Tue salan end en an	Arm B: GSK4381562 plus dostarlimab
Treatment arms	Arm C: GSK4381562 plus dostarlimab plus belrestotug
	Arm D: dostarlimab plus belrestotug
Description	An open-label study of GSK4381562 administered as monotherapy and in combination with anticancer agents
Timeline	Study start: Q1 2022
	Data anticipated: 2026+
Key end points	Safety and PK
Clinicaltrials.gov	<u>Link</u>



Oncology belantamab

NCT05714839 - DREAMM-20

Phase	1/11
Patient	Relapsed/refractory multiple myeloma (RRMM)
Subjects	124
T.,	Part 1: belantamab (may switch to belantamab mafodotin in case of PD)
Treatment arms	Part 2: belantamab and Belamaf
Description	An open-lab multicentre, dose escalation and expansion trial to investigate the safety, tolerability and clinical activity of belantamab as monotherapy and in combination with other treatments in participants with multiple myeloma
Timeline	Trial start: Q3 2023
	Data anticipated: 2026+
Key end points	Part 1: Safety and tolerability (including DLTs), PK and recommended Part 2 dose
	Part 2: Safety and tolerability, PK and recommended phase II dose
Clinicaltrials.gov	<u>Link</u>



Oncology GSK4524101

NCT06077877

Phase	1/11
Patient	Adult participants with solid tumors
Subjects	135
	Arm A, Part 1: GSK4524101 monotherapy
	Arm B, Part 1: GSK4524101 plus niraparib
Treatment arms	Arm C, Part 1: GSK4524101 food effect cohort
	Arm D, Part 2: GSK4524101 plus niraparib
	Arm E, Part 2: Niraparib
Description	A first-time-in-human, open-label, multicentre, dose escalation and expansion study of the oral DNA Polymerase Theta inhibitor (POLQi) GSK4524101 and the PARP inhibitor (PARPi) niraparib in adult participants with solid tumors
Timeline	Trial start: Q4 2023
	Data anticipated: 2026+
Key end points	DLTs, AEs, SAEs, ORR
Clinicaltrials.gov	<u>Link</u>



GSK5733584

NCT06431594

Phase	I
Patient	Adult participants with solid tumors
Subjects	240
Treatment arms	Part 1: Dose escalation with GSK5733584 Part 2: Dose expansion with GSK5733584
Description	A trial to evaluate the safety, tolerability, pharmacokinetics and clinical activity of GSK5733584 for injection in subjects with advanced solid tumors
Timeline	Trial start: 2Q 2024 Data anticipated: 2026+
Key end points	Part 1: DLT Part 2: ORR
Clinicaltrials.gov	<u>Link</u>





linerixibat

NCT04950127 - GLISTEN

Phase	III
Patient	Participants with primary biliary cholangitis (PBC)
Subjects	238
Treatment arms	Arm A: linerixibat Arm B: linerixibat followed by placebo Arm C: placebo Arm D: placebo followed by linerixibat
Description	A two-part randomised, placebo controlled, double blind, multicentre trial to evaluate the efficacy and safety of linerixibat for the treatment of cholestatic pruritus in participants with primary biliary cholangitis
Time aline	Trial start: Q3 2021
Timeline	Data anticipated: H2 2024
Key end points	Change from baseline in monthly itch scores over 24 weeks using Numerical Rating Scale (NRS)
Clinicaltrials.gov	<u>Link</u>



GSK4532990 (NASH/MASH)

NCT05583344 - HORIZON

NCT06104319 - SKYLINE

Phase	IIb	Phase	lla
Patient	Adults with non-alcoholic steatohepatitis (NASH) and advanced fibrosis	Patient	Adult participants with NASH or suspected NASH
Subjects	246	Subjects	48
	Arm 1: high dose GSK4532990	Treatment arms	Arm 1: GSK4532990 Dose 1
Treatment arms	Arm 2: low dose GSK4532990		Arm 2: GSK4532990 Dose 2
	Arm 3: placebo		Arm 3: GSK4532990 Dose 3
Description	A placebo-controlled trial to evaluate the efficacy and safety of GSK4532990		Arm 4: GSK4532990 Dose 4
Description	in adults with advanced non-alcoholic steatohepatitis (NASH)	Description	A single dose, open-label, dose exploration study to assess the PK-PD activity, safety, and tolerability of GSK4532990 in adult participants with NASH or suspected NASH
Timeline	Trial start: Q1 2023		
	Data anticipated: 2025		<u> ' </u>
Key end points	Part 1: Percentage of participants achieving ≥ 1 stage improvement in	Timeline	Trial start: Q1 2024
	histological fibrosis with no worsening of NASH (at week 52)		Data anticipated: 2025
	Part 2: Percentage of participants achieving NASH resolution with no worsening of fibrosis (at week 52)	Key end points	Predicted percent change from baseline in liver biopsy-derived HSD17B13 protein expression levels and mRNA expression levels
Clinicaltrials.gov	<u>Link</u>	Clinicaltrials.gov	<u>Link</u>



GSK4172239 (Sickle cell disease)

NCT05660265

Phase	I .
Patient	Participants with sickle cell disease
Subjects	40
	Cohort 1: GSK4172239D (Dose 1)
	Cohort 2: GSK4172239D (Dose 2)
Treatment arms	Cohort 3: GSK4172239D (Dose 3)
reatment arms	Cohort 4: GSK4172239D (Dose 4)
	Cohort 5: GSK4172239D (Dose 5)
	Food effect cohort
Description	A randomised, placebo-controlled, double-blind (sponsor unblind), parallel group, single dose, dose escalation to evaluate the safety, tolerability and pharmacokinetics of GSK4172239D
Timeline	Trial start: Q3 2023
	Data anticipated: 2025
Key end points	Area under curve zero to time infinity (AUC 0-inf) for GSK4106401 after a single oral dose of GSK4172239D
Clinicaltrials.gov	<u>Link</u>



Glossary



Glossary

ADC	Antibody drug conjugate
AE	Adverse event
AESI	Adverse event of special interest
AIR	At increased risk
AUC	Area under curve
ВСМА	B-cell maturation antigen
BICR	Blinded Independent Central Review
BRCA	Breast cancer
CAE	Corneal adverse events
CBR	Clinical benefit rate
cCR	Complete clinical response
CKD	Chronic kidney disease
CfB	Change from baseline
CMV	Cytomegalovirus
CN	China
COPD	Chronic obstructive pulmonary disease
СР	Cholestatic pruritus
CRR	Complete response rate
CRSwNP	Chronic rhinosinusitis with nasal polyps
cUTI	Complicated urinary tract infection
CV	Cardiovascular
DDI	Drug-drug interaction
DFS	Disease-freee survival
DL	Dose level
DLT	Dose-limiting toxicity
dMMR	Deficient mismatch repair
DoR	Duration of response
DPNP	Diabetic peripheral neuropathic pain
EASI	Eczema Area and Severity Index

EGPA	Eosinophilic granulomatosis with polyangiitis
FVC	Forced vital capacity
GC	Urogenital gonorrhea
GMMA	Generalised Modules for Membrane Antigens
GSI	Gamma secretase inhibitor
HA	
	Healthy adults
HBV	Hepatitis B virus
HES	Hypereosinophilic syndrome
Hgb	Hemoglobin
hSBA	Human serum bactericidal assay
HZ	Herpes zoster
IC	Immunocompromised
ICR	Independent central review
iNTS	Invasive non-typhoidal salmonella
ITT	Intention-to-treat
JP	Japan
LLOQ	Lower limit of quantitation
LRTS	Lower respiratory tract symptoms
MAD	Multiple ascending dose
MAE	Medical attended events
MDI	Metered dose inhaler
MAPS	Mulitple Antigen Presenting System
MASH	Metabolic dysfunction-associated steatohepatit
MM	Multiple myeloma
MMR	Measles, mumps and rubella
MMRV	Measles, mumps, rubella and varicella
MRD	Multiple rising dose
MSI-H	Microsatellite instability high
NASH	Non-alcoholic steatohepatitis
NRS	Numeria Patina Caala
1413	Numeric Rating Scale

NSCLC	Non-small cell lung cancer
OMV	Outer membrane vesicle
ORR	Overall response rate
OS	Overall surival
PBC	Primary biliry cholangitis
PFS	Progression-free survival
PFS2	Time to second disease progression or death
PK	Pharmacokinetic
PMF	Primary myelofibrosis
Post-PV/ET MF	Post-essential thrombocythemia myelofibrosis
RCC	Refractory chronic cough
RL	Repeat dose level
RRMM	Relapsed/refractory multiple myeloma
RSV	Respiratory syncytial virus
SAD	Single ascending dose
SAE	Serious adverse event
siRNA	Small interfering RNA
SoC	Standard of care
SSc-ILD	Systemic sclerosis associated interstitial lung disease
TOC	Test of cure
TTBR	Time to best response
TTD	Time to treatment discontinuation
TTP	Time to tumour progression
TTR	Time to treatment response
UTI	Urinary tract infection
uUTI	Uncomplicated urinary tract infection
VGPR	Very good partial remission
VSP	Vital sign parameters
YoA	Years of age

