Sno	REC Reference	IRAS	Name of Trial	Target Number of Patients Agreed?	Minimum Number of Patients Agreed	Maximum Number of Patients Agreed	Target Date To Recruit Patients Agreed?	Date Agreed to Recruit Target Number of Patients	Total Number of Patients Recruited at the Agreed Target Date	Date That The Trial Closed to Recruitment	Total Number Of Study Participants Recruited	Reason for Closure of Trial
1	14/ES/0001	141631	Randomised, Double-Blind, Placebo-Controlled, Parallel-Group Study to Assess Cardiovascular Outcomes Following Treatment with Ertugliflozin (MK-8835/PF-04971729) in Participants with Type 2 Diabetes Mellitus and Established Vascular Disease	Number Agreed	10	10	Not Available / Not agreed		0	29/11/2019	3	Recruitment Finished
2	14/WM/0049	145163	An openlabel singlearm multicentre noncontrolled phase 3a trial investigating safety and efficacy of nonacog betapegol (N9GP) in prophylaxis and treatment of bleeding episodes in previously untreated patients with haemophilia B (FIX activity ≤2%)	Range Agreed	1	1	Date Agreed	30/09/2019	0	02/03/2020	0	Withdrawn By Host
3	15/LO/0605	161395	Phase I open label, dose escalation trial to determine the MTD, safety, PK and efficacy of afatinib monotherapy inchildren aged 2 years to <18 years with recurrent/refractory neuroectodermal tumours, rhabdomyosarcoma and/orother solid tumours with known ErbB pathway deregulation regardless of tumour histology.	Range Agreed	2	6	Date Agreed	31/12/2019	7	01/06/2020	7	Withdrawn By Sponsor
4	15/LO/1419	183975	A randomized, phase 3 trial with anti-PD-1 monoclonal antibody pembrolizumab (MK-3475) versus placebo for patients with early stage NSCLC after resection and completion of standard adjuvant therapy (PEARLS)	Range Agreed	5	8	Date Agreed	15/11/2019	11	15/11/2019	11	Recruitment Finished
5	16/LO/0086	193561	A Single-Masked, Randomised, Controlled, Parallel Group, Phase 3 Clinical Trial Of Retinal Gene Therapy For Choroideremia Using An Adeno-Associated Viral Vector (AAV2) Encoding Rab Escort Protein 1 (REP1)	Range Agreed	1	10	Date Agreed	30/09/2019	3	02/10/2019	3	Recruitment Finished
6	16/LO/2126	218039	A randomized trial comparing the ELUVIA™ drug- eluting stent versus bare metal self-expanding nitinol stents in the treatment of superficial femoral and/or proximal popliteal arteries.	Range Agreed	1	10	Date Agreed	31/01/2020	1	31/03/2020	1	Recruitment Finished

7	16/NW/0082	184887	Breast reconstruction Outcomes With and without StratticE	Range Agreed	100	200	Date Agreed	31/10/2019	121	31/10/2019	121	Recruitment Finished
8	16/SC/0551	214620	XIRIUS Study (A Dose Escalation, Phase 1/2 Clinical Trial of Retinal Gene Therapy for X- linked Retinitis Pigmentosa Using an Adeno- Associated Viral Vector (AAV8) Encoding Retinitis Pigmentosa GTPase Regulator (RPGR))	Range Agreed	1	15	Date Agreed	28/02/2020	7	20/03/2020	7	Withdrawn By Sponsor
g	17/EM/0121	221773	A randomized, open, multinational, multicentre, 2-part study in spontaneously breathing preterm neonates with mild to moderate respiratory distress syndrome to investigate the safety, tolerability and efficacy of inhaled nebulised poractant alfa (porcine surfactant, curosurf®) in comparison with nCPAP alone	Range Agreed	1	6	Date Agreed	31/05/2022	5	23/04/2020	5	Withdrawn By Sponsor
1	0 17/EM/0122	224376	A Phase 3, Open-Label, Multicenter Study Of ALXN1210 In Children And Adolescents With Atypical Hemolytic-Uremic Syndrome (aHUS)	Range Agreed	1	1	Date Agreed	30/11/2020	0	23/01/2020	0	Recruitment Finished
1	17/EM/0152	216307	A Multi-center, Randomised, Double-blind, Placebo-controlled Phase III Trial of the FLT3 Inhibitor Gilteritinib Administered as Maintenance Therapy Following Allogeneic Transplant for Patients with FLT3/ITD AML	Range Agreed	1	3	Date Agreed	31/03/2020	5	15/02/2020	5	Recruitment Finished
1	2 17/LO/0182	213821	A Prospective, Randomized, Multicenter Controlled Trial of CERAMENTâ,,¢ G as Part of Surgical Repair of Open Diaphyseal Tibial Fractures	Range Agreed	12	20	Date Agreed	30/12/2020	8	15/06/2020	8	Withdrawn By Sponsor
1	3 17/LO/0236	220765	PROMINENT PEMAFIBRATE TO REDUCE CARDIOVASCULAR OUTCOMES BY REDUCING TRIGLYCERIDES IN PATIENTS WITH DIABETES	Range Agreed	5	10	Date Agreed	31/03/2020	6	14/02/2020	6	Recruitment Finished

14	17/LO/0243	219613	A randomized, double-blind, multi-dose, placebo- controlled study to evaluate the efficacy, safety and tolerability ofGSK2330672 administration for the treatment of pruritus in patients with primary biliary cholangitis.(GLIMMER:GSK2330672 triaL of Ibat inhibition with Multidose Measurement for Evaluation of Response).	Range Agreed	1	4	Date Agreed	24/09/2019	0	24/09/2019	0	Recruitment Finished
15	17/LO/0506	220370	A Single Arm, Open-Label, Multi-Centre, Phase I/II Study Evaluating the Safety and Clinical Activity of AUTO3, a CAR T Cell Treatment Targeting CD19 and CD22 in Paediatric and Young Adult Patients with Relapsed Refractory B- cell Acute Lymphoblastic Leukaemia	Range Agreed	10	15	Date Agreed	31/12/2019	11	18/05/2020	11	Withdrawn By Sponsor
16	17/LO/1088	228921	A Phase 3, 2-Arm, Open-label Study to Evaluate the Safety and Pharmacodynamics of Long- termIvacaftor Treatment in Subjects With Cystic Fibrosis Who Are Less Than 24 Months of Age at Treatment Initiation and Have a CFTR Gating Mutation	Range Agreed	1	3	Date Agreed	11/12/2019	0	11/12/2019	0	Recruitment Finished
17	17/LO/1306	228268	A phase III randomized open-label multi-center study of ruxolitinib vs. best available therapy in patients with corticosteroid-refractory chronic graft vs host disease after allogenic stem cell transplantation (REACH 3)	Range Agreed	1	3	Date Agreed	11/10/2019	4	06/12/2019	4	Recruitment Finished
18	17/NE/0200	220486	Strategic MAnagement to Optimize Response To CardiacResynchronization Therapy Registry (SMART Registry)	Number Agreed	12	12	Date Agreed	31/08/2019	27	19/08/2019	27	Recruitment Finished
19	17/NE/0358	220871	Relative bioavailability and comparative pharmacokinetics of 13-CRA oral liquid and extracted capsule formulations: a randomised, open label, multi-dose, cross-over clinical trial in patients requiring treatment cycles of 13-CRA.	Range Agreed	2	3	Date Agreed	30/11/2019	2	17/07/2019	2	Recruitment Finished

20	17/NS/0106	230907	The hydrus microstent for refractory open-angle glaucoma: A prospective, multicenter clinical trial	Range Agreed	3	10	Date Agreed	31/05/2019	3	23/10/2019	3	Recruitment Finished
21	17/NW/0247	222303	A Phase 3, Multicenter, Randomized, Double- Blind, Placebo-Controlled Study to Evaluate the Efficacy	Number Agreed	5	5	Date Agreed	31/03/2025	1	30/09/2019	1	Recruitment Finished
22	17/NW/0399	224760	Study of the measurement of volatile agents to diagnose urological disease	Range Agreed	90	150	Date Agreed	31/07/2019	82	31/07/2019	82	Recruitment Finished
23	17/SC/0245	219967	A Prospective, Open-label, Long-term Safety and Efficacy Study of Teduglutide in Pediatric Patients with Short Bowel Syndrome Who Completed TED-C14-006 or SHP633-301	Range Agreed	1	1	Not Available / Not agreed		0	30/06/2020	2	Recruitment Finished
24	17/SC/0554	225522	A Phase III, multicenter, randomized controlled study to compare safety and efficacy of a haploidentical HSCT and adjunctive treatment with ATIR101, a T-lymphocyte enriched leukocyte preparation depleted ex vivo of host alloreactive T-cells, versus a haploidentical HSCT with post-transplant cyclophosphamide in patients with a hematologic malignancy (HATCY study)	Number Agreed	4	4	Date Agreed	30/12/2020	4	12/11/2019	4	Withdrawn By Sponsor
25	17/WA/0347	234208	A randomised, double-blind, placebo-controlled, parallel-group, multicentre study to demonstrate the effects ofSotagliflozin on Cardiovascular and Renal Events in Patients with Type 2 Diabetes, Cardiovascular Risk Factors andModerately Impaired Renal Function	Number Agreed	12	12	Date Agreed	31/12/2019	12	16/09/2019	12	Recruitment Finished
26	17/WM/030 8	226910	Randomized, Double-Blind, Phase 3B Trial to Evaluate the Safety and Efficacy of 2 Treatment Regimens of Aztreonam 75 mg Powder and Solvent for Nebulizer Solution / Aztreonam for Inhalation Solution (AZLI) in Pediatric Subjects with Cystic Fibrosis (CF) and New Onset Respiratory Tract Pseudomonas aeruginosa (PA) Infection/Colonization	Range Agreed	1	2	Date Agreed	01/11/2019	2	17/02/2020	2	Recruitment Finished

27	17/WM/031 6	226628	A Post-Approval Registry of the TREO [®] Stent- Graft for Patients with Infrarenal Abdominal Aortic Aneurysms	Range Agreed	1	10	Date Agreed	01/06/2020	4	31/12/2019	4	Recruitment Finished
28	17/YH/0426	231118	A Randomized, Double-Blind, Placebo-Controlled Study to Investigate the Efficacy and Safety of Dupilumab Administered Concomitantly with Topical Corticosteroids in Patients, ≥6 Years to <12 Years Of Age, with Severe Atopic Dermatitis	Range Agreed	2	20	Date Agreed	30/09/2018	5	21/08/2019	5	Recruitment Finished
29	18/EE/0005	232671	A Multicenter, Open-Label Study To Estimate The Effect Sizes Of HRCT Endpoints In Response To Glucocorticoid Induction Therapy In Subject With Pulmonary Sarcoidsis	Number Agreed	2	2	Date Agreed	31/08/2019	0	04/09/2019	0	Withdrawn By Sponsor
30	18/EE/0236	242210	Efficacy and safety of Fixed-Dose Combination (FDC) products containing trazodone andgabapentin in patients affected by painful diabetic neuropathy: randomized, controlled, dosefinding study	Range Agreed	1	8	Date Agreed	30/03/2020	1	14/04/2020	1	Recruitment Finished
3:	18/EM/0153	240773	A PHASE 3, RANDOMIZED, DOUBLE- BLIND,PLACEBO-CONTROLLED STUDY TO EVALUATE THEEFFICACY AND SAFETY OF AG- 348 IN NOTREGULARLY TRANSFUSED ADULT SUBJECTS WITHPYRUVATE KINASE DEFICIENCY	Range Agreed	1	2	Date Agreed	28/02/2020	0	05/02/2020	0	Recruitment Finished
32	18/EM/0166	246673	PULMOCIDE - A double-blind, placebo- controlled study to assess the effects of inhaled PC945 in the treatment of culture-positive Aspergillus fumigatus infection in subjects with moderate to severe asthma	Range Agreed	2	3	Date Agreed	31/12/2019	1	01/06/2020	1	Withdrawn By Sponsor
33	18/EM/0365	252018	An open-label extension trial of the long-term safety of nintedanib in patients with Progressive Fibrosing Interstitial Lung Disease (PF-ILD	Number Agreed	3	3	Date Agreed	31/07/2019	3	31/07/2019	3	Recruitment Finished
34	18/ES/0124	248009	An open-label- study to assess the safety, pharmacokinetics and pharmacodynamics of inhaled PC945 in adult Cystic Fibrosis (CF) patients with persistent pulmonary Aspergillus fumigatus infection.	Range Agreed	1	9	Date Agreed	31/12/2019	4	01/06/2020	4	Withdrawn By Sponsor

З	35	18/LO/0656	240062	Multiple escalating dose study of BAY 1093884 in adults with Haemophilia A or B with or without inhibitors	Number Agreed	1	1	Date Agreed	16/07/2020	1	05/08/2019	1	Withdrawn By Sponsor
3	36	18/LO/0711	241788	A Phase 3, Randomized, Double-Blind, Placebo- Controlled, 12-month Study to Evaluate the Efficacy and Safety of MK-7264 in Adult Participants with Chronic Cough (PN030)	Number Agreed	8	8	Date Agreed	07/08/2019	5	12/08/2019	5	Recruitment Finished
3	37	18/LO/0755	245151	A Randomized, Open-label, 24-Week Safety, Efficacy, and Pharmacokinetic Study of Teduglutide in Infants 4 to 12 Months of Age with Short Bowel Syndrome Who are Dependent on Parenteral Support	Range Agreed	1	2	Date Agreed	30/11/2019	1	02/03/2020	2	Recruitment Finished
3	38	18/LO/1007	242697	Patient-Reported Outcomes with the Accu- Chek® SoloMicropump System vs. Insulet OmniPod® vs. Multiple DailyInjection Therapy in Type 1 Diabetes	Range Agreed	1	4	Date Agreed	31/07/2019	4	31/07/2019	4	Recruitment Finished
З	39	18/LO/1311	248599	A phase III, multicenter, randomized, double- masked, active comparatorcontrolled study to evaluate the efficacy and safety of RO6867461 in patients with diabetic macular edema (RHINE)	Number Agreed	2	2	Date Agreed	31/12/2019	2	22/08/2019	2	Recruitment Finished
4	10	18/LO/1370	249657	PREcision Event Monitoring of PatienTs with Heart Failure usingHeartLogicTM	Number Agreed	10	10	Date Agreed	30/12/2023	18	26/06/2020	18	Recruitment Finished
4	11	18/LO/1922	251251	A Phase 2a, Open-label, Single and Multiple Dose Study to Evaluate the Pharmacokinetics, Safety, Tolerability and Treatment Effect of GBT440 in Pediatric Participants with Sickle Cell Disease	Range Agreed	1	3	Date Agreed	31/05/2021	2	06/01/2020	2	Withdrawn By Sponsor
4	12	18/LO/2136	251414	A randomized, multicentre, double-blind, placebo-controlled, phase-III clinical study to evaluate the efficacy and safety of intrathecally administered RO7234292 (RG6042) in patients with manifest Huntington's Disease.	Range Agreed	1	8	Date Agreed	30/06/2020	11	17/04/2020	11	Recruitment Finished

43	18/NE/0102	240702	A phase I, open–label, randomized, pharmacokinetic, pharmacodynamic, and safety study ofEtrolizumab followed by open–label extension and safety monitoring in paediatric patientsFrom 4 years to less than 18 years of age with moderate to severe ulcerative colitis orModerate to severe crohn's disease	Number Agreed	1	1	Date Agreed	31/12/2019	2	20/08/2019	2	Recruitment Finished
44	18/NE/0142	242919	A Phase 3, Double-Blind, Randomized, Placebo- Controlled, Multicenter Study to Evaluate the Efficacy and Safety of Obeticholic Acid in Subjects with Compensated Cirrhosis due to Nonalcoholic Steatohepatitis Protocol 747-304	Range Agreed	1	8	Date Agreed	08/11/2019	1	22/11/2019	1	Recruitment Finished
45	18/NE/0169	243977	A 6-month randomised, double-blind, placebo controlled multicentre parallel group study to evaluate efficacy and safety of bumetanide 0.5mg twice a day followed by an open label active 6-month treatment period with bumetanide (0.5mg twice a day) and a 6 weeks discontinuation period after treatment stop	Range Agreed	6	12	Date Agreed	31/03/2020	11	21/02/2020	11	Recruitment Finished
46	18/NI/0178	253727	A Phase 1 / 2, Drug-Drug Interaction Study of FDL169 and FDL176 in Healthy Subjects and in Cystic Fibrosis Subjects Homozygous for the F508del-CFTR mutation	Range Agreed	1	4	Date Agreed	29/02/2020	0	10/12/2019	0	Withdrawn By Sponsor

47	18/NW/0098	242717	Two-part, double-blind, placebo-controlled, randomized, parallel-group study: (Part 1) in healthy male subjects to assess safety and tolerability of ascending repeated oral doses of BAY 1902607 including its effect on the pharmacokinetics of a sub-therapeutic dose of midazolam (MDZ), followed by (Part 2) a two- way crossover administration of four different doses of BAY 1902607 in patients with refractory chronic cough to assess safety, tolerability and efficacy for proof of concept	Number Agreed	2	2	Date Agreed	30/04/2019	2	05/07/2019	3	Recruitment Finished
48	18/NW/0215	239446	A Phase 2, Randomized, Placebo-Controlled Study Of Safety And Efficacy Following Repeat- Dose Administration Of Evinacumab (Anti- Angptl3) In Patients With Severe Hypertriglyceridemia (Shtg) At Risk For Acute Pancreatitis	Range Agreed	1	3	Date Agreed	30/11/2019	4	09/07/2019	4	Recruitment Finished
49	18/NW/0325	244842	A Randomized, Double-Blind, Placebo-Controlled Study of the Safety and Efficacy of Varying Doses and Dose Regimens of Evinacumab in Patients with Persistent Hypercholesterolemia Despite Maximally Tolerated Lipid Modifying Therapy	Range Agreed	1	3	Date Agreed	31/12/2019	3	24/11/2019	3	Recruitment Finished
50	18/NW/0565	246158	A Phase II randomised , double blind, placebo- controlled, parallel group, multicentre study to evaluate the safety and efficacy of repeated oral doses of Blautix in adult subjects with irritable bowel syndrome (IBS) subtypes IBS-C and IBS-D	Range Agreed	6	14	Date Agreed	31/03/2020	4	17/01/2020	4	Withdrawn By Host
51	18/NW/0599	242748	Study of TRE Seals on Early Post-operative Subjects (STEPS)	Range Agreed	1	10	Date Agreed	29/07/2019	3	30/10/2019	4	Recruitment Finished
52	18/NW/0742	236247	A Phase 2, Randomized, Double-Blind, Placebo- Controlled Efficacy and Safety Study of Palovarotene in Subjects with Multiple Osteochondromas	Range Agreed	4	15	Date Agreed	30/06/2019	3	05/12/2019	7	Withdrawn By Host

53	18/SC/0395	245499	A Multicenter, Randomized, Phase III Registration Trial of Transplantation of NiCord®, Ex Vivo Expanded, Umbilical Cord Blood-derived, Stem and Progenitor Cells, versus Unmanipulated Umbilical Cord Blood for Patients with Hematological Malignancies	Range Agreed	1	3	Date Agreed	31/12/2019	1	24/12/2019	1	Recruitment Finished
54	18/SS/0140	252391	Epidemiological analysis for Hereditary Angioedema Disease (EHA Study)	Number Agreed	100	100	Date Agreed	30/06/2022	11	28/02/2020	11	Withdrawn By Sponsor
55	18/WM/002 2	237623	AN OPEN-LABEL STUDY TO EVALUATE THEEFFICACY AND SAFETY OF AG-348 IN REGULARLYTRANSFUSED ADULT SUBJECTS WITH PYRUVATE KINASE (PK) DEFICIENCY	Range Agreed	1	2	Date Agreed	31/12/2019	1	31/12/2019	1	Recruitment Finished
56	18/WM/031 2	228906	ACCEPTABILITY STUDY ON KETOGENIC DIET SUPPLEMENT USING CAMBROOKE KETOVIE™ MEDICAL FOODS	Range Agreed	1	15	Date Agreed	30/09/2019	11	30/09/2019	11	Recruitment Finished
57	18/YH/0055	241341	A Phase 2, Prospective, Randomized, Open-label Study on the Efficacy of Defibrotide Added to Standard of Care Immunoprophylaxis for the Prevention of Acute Graft-versus- Host-Disease in Adult and Pediatric Patients After Allogeneic Hematopoietic Stem Cell Transplant	Range Agreed	1	2	Date Agreed	31/12/2019	4	31/10/2019	4	Recruitment Finished
58	18/YH/0136	214921	A multi-centre, pilot, prospective, trial of DermaRep [™] Device in the treatment of venous leg ulcers	Range Agreed	1	5	Date Agreed	30/09/2019	6	30/09/2019	6	Recruitment Finished
59	19/EE/0091	256412	An open-label, pilot study to assess safety, tolerability, pharmacokinetics and effects of inhaled PC945 in the pre-emptive treatment of Aspergillus fumigatus colonisation in lung transplant recipients	Range Agreed	1	4	Date Agreed	28/02/2020	0	01/06/2020	0	Withdrawn By Sponsor
60	19/EM/0003	251874	A Randomized, Double-Blind, Placebo Controlled, Global Phase 3 Study Of Edasalonexent In Pediatric Patients With Duchenne Muscular Dystrophy	Range Agreed	1	3	Date Agreed	30/09/2019	5	30/09/2019	5	Recruitment Finished

61	19/EM/0047	257111	A Phase III, Multicenter, Randomized, Double- Masked, Active Comparator-Controlled Study To Evaluate The Efficacy And Safety Of Faricimab In Patients With Neovascular Age-Related Macular Degeneration (TENAYA)	Number Agreed	4	4	Date Agreed	31/12/2019	3	24/10/2019	3	Recruitment Finished
62	19/EM/0094	259038	Effects of N-Acetyl-L-Leucine on Niemann-Pick disease type C: A multinational, multicenter, open-label, rater-blinded Phase II study.	Range Agreed	1	4	Date Agreed	31/01/2020	1	31/01/2020	1	Recruitment Finished
63	19/EM/0220	265213	A Phase III, randomized, multicenter, open-label, non-inferiority studyevaluating the efficacy, safety and tolerability of switching to dolutegravir/lamivudine fixed dose combination in HIV-1 infected adults who are virologically suppressed	Range Agreed	1	2	Date Agreed	31/05/2020	0	20/04/2020	0	Withdrawn By Sponsor
64	19/LO/0157	256597	A study to evaluate safety, reactogenicity and immunogenicity of GSK Biologicals' RSV investigational vaccine based on viral proteins encoded by chimpanzee-derived adenovector (ChAd155-RSV) (GSK3389245A) in infants	Number Agreed	2	2	Date Agreed	30/09/2019	1	20/10/2019	1	Recruitment Finished
65	19/LO/0351	256743	A Phase 3, Randomized, Double-Blind, Study Comparing Risankizumab to Placebo in Subjects with Active Psoriatic Arthritis (PsA) Who Have a History of Inadequate Response to or Intolerance to at Least One Disease Modifying Anti- Rheumatic Drug (DMARD) Therapy	Range Agreed	1	3	Date Agreed	30/06/2020	0	11/03/2020	0	Recruitment Finished
66	19/LO/0352	257478	A Phase 3, Randomized, Double-Blind Study Comparing Risankizumab to Placebo in Subjects with Active Psoriatic Arthritis Including Those Who Have a History of Inadequate Response or intolerance to Biologic Therapy(ies)	Range Agreed	1	3	Date Agreed	30/06/2020	0	28/11/2019	0	Withdrawn By Sponsor

67	19/LO/1317	268821	A Phase 3, Open-label Study Evaluating the Long- term Safety and Efficacy of VX-445 Combination Therapy in Subjects With Cystic Fibrosis Who Are Heterozygous for the F508del Mutation and a Gating or Residual Function Mutation (F/G and F/RF Genotypes)	Range Agreed	1	3	Date Agreed	30/09/2020	2	26/05/2020	2	Recruitment Finished
68	19/LO/1318	268820	A Phase 3, Randomized, Double-blind, Controlled Study Evaluating the Efficacy and Safety of VX-445 Combination Therapy in Subjects With Cystic Fibrosis Who Are Heterozygous for the F508del Mutation and a Gating or Residual Function Mutation (F/G and F/RF Genotypes)	Range Agreed	1	3	Date Agreed	31/07/2020	2	26/05/2020	2	Recruitment Finished
69	19/NW/0026	249432	VX18-121-101: A Phase 2, Randomized, Double- blind, Controlled Study to Evaluate the Safety and Efficacy of VX-121 Combination Therapy in Subjects Aged 18 Years and Older with Cystic Fibrosis	Range Agreed	1	4	Date Agreed	30/09/2019	6	30/08/2019	6	Recruitment Finished
70	19/NW/0181	260431	A Randomised, Double-blind, Placebo- Controlled, Crossover, Dose Escalation Study of BLU-5937 in Subjects with Unexplained or Refractory Chronic Cough (RELIEF)	Range Agreed	1	5	Date Agreed	31/01/2020	5	04/04/2020	6	Withdrawn By Sponsor
71	19/NW/0482	269475	A Phase 3b, Randomized, Double-blind, Controlled Study Evaluating the Efficacy and Safety of VX-445/Tezacaftor/Ivacaftor in Cystic Fibrosis Subjects, Homozygous for F508del	Range Agreed	1	5	Date Agreed	30/03/2020	5	19/12/2019	5	Withdrawn By Sponsor
72	19/NW/0540	269907	A Phase 3, Open-label Study Evaluating the Long- term Safety of VX-445 Combination Therapy in Subjects With Cystic Fibrosis	Range Agreed	1	3	Date Agreed	31/07/2020	4	29/05/2020	4	Recruitment Finished
73	19/SC/0031	257722	A Randomized, Placebo-controlled, Double- blind, Phase 1/2a Study to Evaluate the Safety, Reactogenicity, and Immunogenicity of Ad26.RSV.preF in RSV-seronegative Toddlers 12 to 24 Months of Age	Range Agreed	1	5	Date Agreed	30/11/2019	0	06/08/2019	0	Recruitment Finished

74	19/YH/0093	255567	ATLAS - A Randomized Open-label, Phase 1b Study of the Safety of Pirfenidone Solution for inhalation (AP01) in Patients with Idiopathic Pulmonary Fibrosis	Range Agreed	3	5	Date Agreed	16/03/2020	7	29/04/2020	7	Recruitment Finished
75	20/NE/0104	282007	A Phase 3 Randomized Study to Evaluate the Safety and Antiviral Activity of Remdesivir (GS- 5734™) in Participants with Severe COVID-19	Range Agreed	1	9	Date Agreed	31/05/2020	1	29/05/2020	1	Recruitment Finished
76	20/NW/0168	281317	A randomised double-blind placebo-controlled trial to determine the safety and efficacy of inhaled SNG001 (IFN-β1a for nebulisation) for the treatment of patients with confirmed SARS- CoV-2 infection	Range Agreed	8	20	Date Agreed	15/05/2021	8	11/05/2020	8	Recruitment Finished