2 15/W 3 18/L			Name of Trial	Of Patients Agreed?	Number Of Patients Agreed	Number Of Patients Agreed	To Recruit Patients Agreed?	to recruit target number of patients	Patients Recruited At The Agreed Target Date	The Trial Closed To Recruitment	Number Of Study Participants Recruited	Reason For Closure Of Trial
2 15/W 3 18/L	1000100	164000	A dark Deserves		20	20	Data Armad	24/04/2010	10	24/04/2010	10	Recruitment
<u>3 18/L</u>	/NW/0142	164099	Adapt Response An Open Label, Randomised, Pre-surgical, Pharmacodynamics Study to	Number Agreed	20	20	Date Agreed	31/01/2019	10	31/01/2019	10	Finished
<u>3 18/L</u>	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	172946	Compare the Biological Effects of AZD9496 versus Fulvestrant in Postmenopausal Women with ER positive HER-2 negative Primary Breast Cancer	Number Agreed	1	1	Date Agreed	31/12/2018	6	11/01/2019	6	Recruitment Finished
	VVIVI/0455	172940	A Randomized, Double Masked, Uncontrolled, Multicenter Phase I/II Study	Number Agreeu	1	1	Date Agreeu	51/12/2018	0	11/01/2019	0	Fillistieu
4 18/E	/LO/0782	245936	to Evaluate Safety and Tolerability of PAN-90806 Eye Drops, Suspension in Treatment-Na?ve Participants with Neovascular Age-Related Macular Degeneration (AMD)	Range Agreed	1	4	Date Agreed	01/03/2019	1	01/03/2019	1	Recruitment Finished
	/EM/0228	248988	A double blind, placebo-controlled study to assess the anti-viral effect, safety and tolerability of inhaled PC786 for the treatment of acute respiratory syncytial virus (RSV) infection in adult hematopoietic stem cell transplant recipients	Number Agreed	3	3	Date Agreed	11/03/2019	1	12/03/2019	1	Recruitment Finished
			External Natural History Controlled, Open-Label Intervention Study to Assess									
	(the Efficacy and Safety of Long-Term Treatment with Raxone? in Leber?s			_		/ /	_		_	Recruitment
5 16/1	/NE/0255	200900	Hereditary Optic Neuropathy (LHON)	Range Agreed	1	3	Date Agreed	08/03/2019	7	08/03/2019	7	Finished
6 17/L	/LO/0113	219400	A Phase 3, Double-blind, Randomized, Placebo-controlled, Multicenter Study of GBT440 Administered Orally toPatients With Sickle Cell Disease	Range Agreed	1	6	Date Agreed	30/06/2018	2	19/03/2019	2	Recruitment Finished
7 17/9	150/0227	220062	A multi-centre, double-blind, parallel-group, randomised, placebo controlled phase II a study to investigate safety, tolerability, pharmacodynamics, and pharmacokinetics of different doses of orally administered BI 1467335 during a12 work treatment as	Number Agreed	4	4	Data Agroad	19/12/2019	1	07/02/2010	1	Recruitment
/ 1//3	/SC/0237	220963	during a12-week treatment pe	Number Agreed	4	4	Date Agreed	18/12/2018	1	07/02/2019	1	Finished
8 16/L	/LO/0878	201070	Prospective, multi-centre study to evaluate the everlinQ endoAVF System when used to create an endovascular arteriovenous fistula (endoAVF) for patients who require vascular access for haemodialysis.	Range Agreed	15	20	Date Agreed	31/12/2019	13	06/03/2019	13	Recruitment Finished
9 15/E	/EM/0543	176792	A Randomized, Open-label, Controlled Phase 3 Trial to Investigate the Efficacy, Safety, and Tolerability of the BiTE Antibody Blinatumomab as Consolidation Therapy Versus Conventional Consolidation Chemotherapy in Pediatric Subjects With High-risk Fi	Range Agreed	1	5	Date Agreed	30/11/2018	1	11/03/2019	1	Withdrawn By Host
			A Phase 3, Double-Blind, Randomized, Long-Term, Placebo-Controlled,									
	6 <i>1</i>		Multicenter Study Evaluating the Safety and Efficacy of Obeticholic Acid in					/ /				Recruitment
10 15/Y	/YH/0478	186697	Subjects with Nonalcoholic Steatohepatitis	Range Agreed	1	4	Date Agreed	30/06/2019	4	17/06/2019	4	Finished
11 16/5	/SC/0615	210762	A Phase 4 open-label randomized controlled study COmparing the effectiveness of adalimumab iNTROduction andmethotrexate dose escaLation in subjects with Psoriatic Arthritis (CONTROL)	Range Agreed	1	5	Date Agreed	15/10/2018	11	07/05/2019	12	Recruitment Finished
12 16/W	120/0012	210,02										

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
13	16/WM/0511	218115	A Phase 3, Prospective, Multicenter, Uncontrolled, Open-Label Clinical Study to Determine the Efficacy, Safety, and Tolerability of rVWF with or without ADVATE in the Treatment and Control of Bleeding Episodes, the Efficacy and Safety of rVWF in Elec	Number Agreed	1	1	Date Agreed	31/12/2019	0	23/04/2019	0	Recruitment Finished
14	17/EE/0474	229785	An Open-Label, Randomized, Multi-Center, Parallel Group, Two-Arm Study to Assess the Safety, Overall Tolerability, and Antiviral Activity of Brincidofovir versus Standard of Care for Treatment of Adenovirus Infections in High-Risk Pediatric Allogenei	Range Agreed	1	4	Date Agreed	31/05/2019	3	09/05/2019	3	Withdrawn By Sponsor
15	17/EM/0063	213979	A Phase 3 Multicenter, Randomized, Double-Blind, Placebo-Controlled Trial of the FLT3 Inhibitor Gilteritinib (ASP2215) Administered as Maintenance Therapy Following Induction/Consolidation Therapy for Subjects with FLT3/ITD AML in First Complete Remi	Range Agreed	1	3	Date Agreed	31/03/2019	0	10/04/2019	0	Withdrawn By Sponsor
16	17/EM/0404	223875	A Phase 3, Multi-center, Randomized, Double-Masked Study to Evaluate the Clinical Efficacy and Safety of SHP640 (PVP-lodine 0.6% and Dexamethasone 0.1%) Ophthalmic Suspension Compared to PVP-lodine and Placebo in the Treatment of Adenoviral Conjuncti	Range Agreed	1	10	Date Agreed	31/12/2020	2	03/05/2019	2	Withdrawn By Sponsor
17		219676	CardioMEMS Heart Failure System "Outside US" Post-Approval Study	Range Agreed	1	10	Date Agreed		7	21/06/2019	7	Recruitment Finished
18	17/LO/0372	220433	A phase III randomized open-label multi-center study of ruxolitinib vs. best available therapy in patients with corticosteroid-refractory acute graft vs. host disease after allogenic stem cell transplantation A randomized, controlled, open-label, multiple ascending dose study of intravenous brincidofovir in adult allogeneic hematopoietic cell transplant	Range Agreed	1	3	Date Agreed	07/05/2019	3	07/05/2019	3	Recruitment Finished Withdrawn
19		235819	recipients with adenovirus viremia A randomised, double-blind, controlled, parallel-group, multi-country study to investigate the effect of a partially hydrolysed infant formula with added synbiotics on gut microbiota composition and clinical effectiveness in infants	Range Agreed	1	2	Date Agreed		0	09/05/2019	0	By Sponsor Recruitment
20 21	17/NE/0124 17/NI/0096	216591	at high risk of d HOPE-1 (HYDRATION FOR OPTIMAL PULMONARY EFFECTIVENESS)	Range Agreed	2	3	Date Agreed	10/11/2018	2	28/02/2019	1	Finished Withdrawn By Host
	17/NW/0209	213231	An open-label ascending dose cohort study to assess the safety, pharmacokinetics, and preliminary efficacy ofneoGAA (GZ402666) in patients with infantile-onset Pompe disease treated with alglucosidase alfa who demonstrateclinical decline or sub-optim	Range Agreed	1	3	Date Agreed		1	19/04/2019	1	Recruitment Finished
23	17/SC/0462	228889	A Randomized, Double-blind Phase 1/2a Study to Evaluate the Safety, Tolerability and Immunogenicity of Ad26.RSV.preF in Adults 18 to 50 Years of Age and RSV-Seropositive Toddlers 12 to 24 Months of Age	Number Agreed	4	4	Date Agreed	31/03/2018	0	14/06/2019	4	Recruitment Finished

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
24	17/SW/0221	232448	A randomized, partially-blinded, active-controlled,multicenter study of secukinumab to demonstrate reduction of radiographic progression versus GP2017 (adalimumab biosimilar) at 104 weeks and to assess the long term safety, tolerability and efficacy	Number Agreed	4	4	Date Agreed	02/11/2021	0	16/05/2019	0	Withdrawn By Sponsor
	17/3W/0221	249183	A Phase I, Multicenter, Open-Label, Single-Dose, Dose-Ranging Study to Assess the Safety and Tolerability of SB-913, a rAAV2/6-based Gene Transfer in Subjects with Mucopolysaccharidosis II (MPSII)	Range Agreed	1	4	Date Agreed		0	25/04/2019	0	Withdrawn By Sponsor
25		249185	A prospective follow up study to assess performance, safety and efficacy of the PICO 7 NPWT system for surgically closed incision sites and skin grafts	Range Agreed	5	8	Date Agreed		9	21/05/2019	9	Recruitment Finished
27	15/NE/0278	183395	An open-label, randomized, active-controlled,parallel-group, Phase-3b study of theefficacy, safety, and tolerability of 2 mg afliberceptadministered by intravitreal injectionsusing two different treatment regimens to subjects withneovascular age-rela	Number Agreed	5	5	Date Agreed	20/04/2010	9	28/02/2019	9	Recruitment Finished
	14/LO/0122	141557	A multicentre, openlabel, nonrandomised, phase I dose escalation study of regorafenib (BAY 734506) in paediatric subjects with solid malignant tumours that are recurrent or refractory to standard therapy.	Number Agreed	2	2	Date Agreed		3	31/03/2019	3	Recruitment Finished
29	16/LO/0718	203652	A phase 3, multicentre, single-arm, open-label study of the efficacy and safety of B-domain deleted recombinant porcine factor VII (BAX802) in subjects with congenital haemophilia A with factor VIII inhibitors undergoing surgical or other invasive pr	Range Agreed	1	2	Date Agreed	01/07/2020	0	22/05/2019	0	Withdrawn By Host
30	17/LO/1103	230709	Two-part, double-blind, placebo-controlled, randomized, parallel-group study: (Part 1) in healthy male volunteers to assess safety and tolerability of ascending repeated oral doses of BAY 1817080, followed by (Part 2), two- way crossover administratio	Number Agreed	6	6	Date Agreed	26/11/2018	5	06/03/2019	6	Recruitment Finished
31	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
			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-									Recruitment
32	17/NE/0358	220871	CRA. A Phase 3, Multicenter, Randomized, Double-Blind, Placebo-Controlled Study	Range Agreed	2	3	Date Agreed	30/11/2019	2	17/07/2019	2	Finished
33	17/NW/0247	222303	to Evaluate the Efficacy	Number Agreed	5	5	Date Agreed	31/03/2025	1	30/09/2019	1	Recruitment Finished
34	17/NW/0399	224760	Study of the measurement of volatile agents to diagnose urological disease A randomised, double-blind, placebo-controlled, parallel-group, multicentre study to demonstrate the effects of Sotagliflozin on Cardiovascular and Renal	Range Agreed	90	150	Date Agreed	31/07/2019	82	31/08/2019	82	Recruitment Finished
35	17/WA/0347	234208	Events in Patients with Type 2 Diabetes, Cardiovascular Risk Factors andModerately Impaired Renal	Number Agreed	12	12	Date Agreed	31/12/2019	12	16/09/2019	14	Recruitment Finished

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
			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									Recruitment
36	17/YH/0426	231118	Dermatitis	Range Agreed	2	20	Date Agreed	30/09/2018	5	21/08/2019	5	Finished
			A double blind (sponsor open) placebo-controlled, stratified, parallelgroup					- · ·				
			study to evaluate the efficacy and safety of repeat doses of GSK3772847									
			inparticipants with moderate to severe asthma with allergic fungal airway									Withdrawn
37	17/YH/0432	236091	disease (AFAD).	Range Agreed	3	5	Date Agreed	28/06/2019	0	30/06/2019	0	By Sponsor
			A Multicenter, Open-Label Study To Estimate The Effect Sizes Of HRCT									
38	18/EE/0005	232671	Endpoints In Response To Glucocorticoid Induction Therapy In Subject With Pulmonary Sarcoidsis	Number Agreed	2	2	Data Agroad	31/08/2019	0	04/09/2019	0	Withdrawn
50	18/11/0003	232071		Number Agreed	2	2	Date Agreed	51/06/2019	0	04/09/2019	0	By Sponsor Recruitment
39	18/LO/0190	227705	Clinical investigation of the eyeWatch glaucoma drainage device	Number Agreed	15	15	Date Agreed	31/05/2019	15	31/05/2019	15	Finished
			Multiple escalating dose study of BAY 1093884 in adults with Haemophilia A	0			Ŭ					Withdrawn
40	18/LO/0656	240062	or B with or without inhibitors	Number Agreed	1	1	Date Agreed	16/07/2020	1	05/08/2019	1	By Sponsor
			A Phase 3, Randomized, Double-Blind, Placebo-Controlled, 12-month Study									
	40/10/0744	244700	to Evaluate the Efficacy and Safety of MK-7264 in Adult Participants with		2			07/00/2010	-	42/00/2040	-	Recruitment
41	18/LO/0711	241788	Chronic Cough (PN030) A Phase 3, Randomized, Double-Blind, Placebo-Controlled, 12-Month Study	Number Agreed	8	8	Date Agreed	07/08/2019	5	12/08/2019	5	Finished
			to Evaluate the Efficacy and Safety of MK-7264 in Adult Participants with									Recruitment
42	18/LO/0712	241782	Chronic Cough (PN027)	Number Agreed	8	8	Date Agreed	07/08/2019	8	17/05/2019	8	Finished
				0			Ŭ					
			Patient-Reported Outcomes with the Accu-Chek? SoloMicropump System vs.									Recruitment
43	18/LO/1007	242697	Insulet OmniPod? vs. Multiple DailyInjection Therapy in Type 1 Diabetes	Range Agreed	1	4	Date Agreed	31/07/2019	4	31/07/2019	4	Finished
			A phase III, multicenter, randomized, double-masked, active									
44	18/LO/1311	248599	comparatorcontrolled study to evaluate the efficacy and safety of		2	2	Data Agroad	21/12/2010	2	22/08/2010	2	Recruitment Finished
44	16/10/1511	240599	RO6867461 in patients with diabetic macular edema (RHINE)	Number Agreed	2	2	Date Agreed	31/12/2019	2	22/08/2019	2	Fillistieu
			A phase I, open?label, randomized, pharmacokinetic, pharmacodynamic, and									
			safety study of Etrolizumab followed by open? label extension and safety									
			monitoring in paediatric patientsFrom 4 years to less than 18 years of age									Recruitment
45	18/NE/0102	240702	with moderate to severe ulcerat	Number Agreed	1	1	Date Agreed	31/12/2019	2	20/08/2019	2	Finished
			A Phase 3, Open-label Study Evaluating the Long-term Safety and Efficacy of									Recruitment
46	18/NE/0266	241641	VX-659 Combination Therapy in Subjects With Cystic Fibrosis Who Are Homozygous or Heterozygous for the F508del Mutation	Range Agreed	2	3	Date Agreed	31/01/2010	4	20/02/2019	4	Finished
-40	10/11/0200	271041	Two-part, double-blind, placebo-controlled, randomized, parallel-group	Nullec Agi ceu	2	5	Dute Agreeu	51/01/2015	4	20/02/2019	+	rinsheu
			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									Recruitment
47	18/NW/0098	242717	pharmacokinetics of a sub-th	Number Agreed	2	2	Date Agreed	30/04/2019	2	05/07/2019	3	Finished
			A Dhose 2. Devidemined Devide Divid Disaster Controlled Office									\A/i+b al
10	18/NW/0742	236247	A Phase 2, Randomized, Double-Blind, Placebo-Controlled Efficacy and Safety Study of Palovarotene in Subjects with Multiple Osteochondromas		4	15	Date Agreed	30/06/2010	3	09/08/2019	6	Withdrawn By Host
48	10/11/0/42	230247	Study of Palovarotene in Subjects with Multiple Osteochondromas	Range Agreed	4	13	Date Agreed	20/00/2019	5	05/00/2019	0	by HUSL

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10	40/0004/0004	244427	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	Danas Asus d	4	2	Data Associ	24 /00 /2040	2	40/04/2040	2	Recruitment
49	18/WM/0204	244427	Homozygous or Heterozygous for the F508del Mutation	Range Agreed	1	3	Date Agreed	31/08/2019	3	18/04/2019	3	Finished
50	19/EM/0003	251874	A Randomized, Double-Blind, Placebo Controlled, Global Phase 3 Study Of Edasalonexent In Pediatric Patients With Duchenne Muscular Dystrophy VX18-121-101: A Phase 2, Randomized, Double-blind, Controlled Study to	Range Agreed	1	3	Date Agreed	30/09/2019	5	30/09/2019	5	Recruitment Finished
			Evaluate the Safety and Efficacy of VX-121 Combination Therapy in Subjects									Recruitment
51	19/NW/0026	249432	Aged 18 Years and Older with Cystic Fibrosis	Range Agreed	1	4	Date Agreed	30/09/2019	6	30/08/2019	6	Finished
52	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
53	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	10	15/11/2019	10	Recruitment Finished
54	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 Recruitment
55	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	Finished
56	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 Iba	Range Agreed	1	4	Date Agreed	24/09/2019	0	24/09/2019	0	Recruitment Finished
57	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
FO	17/NS/0100	220007	The hydrus microstent for refractory open-angle glaucoma: A prospective,	Paper Agroad	2	10	Data Agreed	21/05/2010	2	22/10/2010	2	Recruitment
58	17/NS/0106	230907	multicenter clinical trial 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	Range Agreed	3	10	Date Agreed		3	23/10/2019	3	Finished Withdrawn
59	17/SC/0554	225522	alloreactive T-cells, versus a	Number Agreed	4	4	Date Agreed	30/12/2020	4	12/11/2019	4	By Sponsor
60	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

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
61	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
62	18/NI/0240	258468	An Observational, Prospective Multicentre Clinical Study to assess the safety and clinical performance of a New Single-use Negative Pressure Wound Therapy System (PICO 7Y) for the Simultaneous Management of Bilateral Closed Incisions in Oncoplastic B	Number Agreed	10	10	Date Agreed	31/12/2019	13	18/05/2019	13	Recruitment Finished
63	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	3	Recruitment Finished
			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									Recruitment
64	18/SC/0395	245499	with Hematological Malignancies AN OPEN-LABEL STUDY TO EVALUATE THEEFFICACY AND SAFETY OF AG-348 IN REGULARLYTRANSFUSED ADULT SUBJECTS WITH PYRUVATE KINASE (PK) DEFICIENCY	Range Agreed Range Agreed	1	3	Date Agreed	31/12/2019	0	24/12/2019	0	Finished Recruitment Finished
	18/WM/0312	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
67	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	Range Agreed	1	2	Date Agreed	31/12/2019	4	31/10/2019	4	Recruitment Finished
68	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
69	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
70	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
			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	Numbe Agreed								Recruitment
71	18/NW/0215	239446	Pancreatitis A Randomized, Double-Blind, Placebo-Controlled Study of the Safety and	Range Agreed	1	3	Date Agreed	30/11/2019	4	09/07/2019	4	Finished
72	18/NW/0325	244842	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