

No	REC Reference Number	IRAS No	Title	Date of First Patient Recruitment	Benchmark Met	Date site invited	Date site selected	HRA Approval date	Date site confirmed by sponsor	Date site confirmed	Date site ready to start	A - Permissions delayed / denied	B - Suspended by Sponsor	C - Closed by Sponsor	D - Sponsor delays	E - Staff Availability issues	F - No Eligible Patients (No patient seen)	G - No Patients Consented	H - Contracting Delays	I - Rare disease	J - Other	Comments	Source of delay			
1	16/YH/0459	213518	Optimal Pathway for Treating neuropathic pain in Diabetes Mellitus (OPTION-DM) trial		N/A	07/10/2016	19/09/2017	24/07/2017	26/09/2017	26/09/2017	02/10/2017												N/A as 70 day benchmark not exceeded at time of report period			
2	16/NW/053	195854	Efficacy and safety of a Double Icodextrin Dose in elderly incident CAPD patients on Incremental peritoneal dialysis therapy: The DiDo Study		N/A	22/06/2016	14/08/2017	02/11/2016	31/08/2017	08/09/2017														N/A as 70 day benchmark not exceeded at time of report period		
3	17/NW/0180	220257	A Phase 2, Open-Label, Single-Arm, Multicenter Study to Evaluate the Efficacy and Safety of INCB054828 in Subjects With Advanced/Metastatic or Surgically Unresectable Cholangiocarcinoma Including FGFR2 Translocations Who Failed Previous Therapy		N/A	06/10/2016	26/07/2017	04/04/2017	26/07/2017	03/08/2017	22/09/2017														N/A as 70 day benchmark not exceeded at time of report period	
4	17/LO/0440	217506	Randomized, Double-Blind, Placebo-Controlled Study to Evaluate the Safety, Tolerability, Pharmacokinetics and Pharmacodynamics of Multiple Ascending Doses of Intrathecally Administered ISIS 814907 in Patients with Mild Alzheimer's Disease		N/A	24/11/2016	27/07/2017	20/06/2017	27/07/2017	03/08/2017	22/08/2017														N/A as 70 day benchmark not exceeded at time of report period	
5	16/NS/0094	206213	Bioimpedance Spectroscopy To Maintain Renal Output: The BISTRO Trial		N/A	05/05/2017	04/08/2017	04/10/2016	11/08/2017	11/08/2017	06/09/2017														N/A as 70 day benchmark not exceeded at time of report period	
6	16/EM/0344	180454	The Effect of MultiPoint Pacing on reverse remodelling and the incidence of ventricular arrhythmias – The MPP-VARR Study		N/A	16/06/2017	31/08/2017	26/10/2016	05/09/2017	05/09/2017	07/09/2017														N/A as 70 day benchmark not exceeded at time of report period	
7	17/NW/0374	226708	A Multi-Centre, Randomised, Open-Label, Controlled Trial Evaluating the Efficacy and Safety of Prophylactic Administration of Concizumab in Haemophilia A and B Patients with Inhibitors		N/A	12/06/2017	29/08/2017	18/07/2017	29/08/2017	05/09/2017	04/10/2017														N/A as 70 day benchmark not exceeded at time of report period	
8	13/LO/0451	105864	The United Kingdom transcatheter aortic valve implantation trial (UK TAVI)		N/A	24/07/2017	16/08/2017	15/06/2017	30/08/2017	08/09/2017	19/09/2017														N/A as 70 day benchmark not exceeded at time of report period	
9	16/SC/0651	213404	An evaluation of the tolerance, compliance and acceptability of a ready to use, liquid, high energy, high protein, peptide-based feed for adults in need of nutrition support – a pilot study		N/A	14/07/2017	26/07/2017	13/01/2017	27/07/2017	03/08/2017	21/08/2017														N/A as 70 day benchmark not exceeded at time of report period	
10	16/LO/1004	207544	Modulation of immune response and outcomes in ALS using IL-2 (MIROCALS-IL2)	11/07/2017	No	09/09/2016	16/03/2017	06/10/2016	21/03/2017	04/04/2017	10/07/2017				Y										Sponsor delayed confirmation of study open to recruitment at site	Sponsor
11	16/NE/0253	185463	Rituximab - a potential cure for the young patient with Graves' disease		No	07/04/2016	05/10/2016	22/09/2016	07/02/2017	17/02/2017	10/05/2017								Y						Contracting delays by Sponsor, further exacerbated by incomplete funding information from Sponsor.	Sponsor

12	16/EE/0370	198596	MesoTRAP: A feasibility study comparing video-assisted thoracoscopic partial pleurectomy/decortication with indwelling pleural catheter in patients with trapped lung and pleural effusion due to malignant pleural mesothelioma designed to address recruitment and randomisation uncertainties and sample size requirements for a phase III trial.	No	16/12/2016	05/05/2017	11/01/2017	05/05/2017	09/05/2017	15/09/2017									Sponsor delayed confirmation of study open to recruitment at site	Sponsor
13	16/NW/0396	191940	Very Low Dose Dexamethasone versus Placebo For the Treatment of Ventilator Dependent Preterm Babies.	No	22/09/2016	24/10/2016	30/06/2016	16/02/2017	22/02/2017	24/07/2017							Y		Contracting delays with sponsor. Sponsor delayed confirmation of study to open at site and delay in provision of IMP.	Sponsor
14	16/YH/0212	200145	Unilateral and Bilateral Neurodynamic Sliding Techniques as a Means of Treating Non-compressive Sciatic Leg Pain: A Pilot Study for a Randomised Controlled Trial	05/01/2017	No	27/06/2016	25/10/2016	25/10/2016	25/10/2016	25/10/2016							Y		Sponsor delay as a result of a protocol amendment - change of venue of intervention.	Sponsor
15	16/LO/0529	182152	CORE: A randomized trial of conventional care versus radioablation (stereotactic body radiotherapy) for extracranial oligometastases	No	20/10/2016	20/10/2016	13/07/2016	31/07/2017	20/07/2017	04/09/2017							Y	Y	Delay by NHS England to list site as an approved contractor to deliver SABR treatment which was required before site can proceed with study.	Neither
16	16/EE/0234	194284	Three versus five years of adjuvant imatinib as treatment of patients with operable GIST with a high risk for recurrence: A randomised phase III study.	No	15/06/2016	25/05/2017	12/09/2016	25/05/2017	25/05/2017	19/07/2017							Y		Sponsor delay in site initiation. Eligible patients seen chose not to participate in study.	Sponsor
17	16/SC/0147	183044	Randomised double-blind cross-over study of a DPP4 inhibitor, SGLT2 inhibitor and thiazolidinedione as third line therapy in patients with type 2 diabetes who have suboptimal glycaemic control on dual therapy with metformin and a sulphonylurea.	22/03/2017	No	08/03/2016	13/12/2016	07/07/2016	14/12/2016	13/01/2017	20/02/2017						Y		Sponsor delayed confirmation of study open to recruitment at site	Sponsor
18	16/YH/0186	200765	A phase 4 double-blind, randomized, placebo-controlled, multi-center study to evaluate the efficacy, safety and tolerability of mirabegron in men with overactive bladder (OAB) symptoms while taking the alpha-blocker tamsulosin hydrochloride for lower urinary tract symptoms (LUTS) due to benign prostatic hyperplasia (BPH).	No	29/04/2016	10/10/2016	11/08/2016	11/10/2016	26/10/2016	29/11/2016							Y	Y	Sponsor delayed confirmation to study open to recruitment at site. Patients screened but no eligible patients identified	Sponsor
19	08/H1002/71	3586	A Pilot Randomised Study to Compare Combination Antibiotic Therapy (Ciprofloxacin, DoxyCycline and Hydroxychloroquine) with standard therapy (Budesonide) in the treatment Of active Crohn's disease.	No	02/02/2016	03/11/2016	14/12/2016	06/06/2017	18/07/2017	03/08/2017							Y	Y	Contracting delays with Sponsor	Sponsor

69	17/LO/0402	209419	A Phase III, multicentre, randomised, placebo-controlled, double-blind study of Atezolizumab (anti-PD-L1 antibody) as adjuvant therapy in patients with PD-L1-selected renal cell carcinoma at intermediate to high risk of developing metastasis following nephrectomy. IMMOTION	25/08/2017	Yes	20/01/2017	30/06/2017	09/06/2017	04/07/2017	07/07/2017	26/07/2017								
70	16/YH/0450	213782	20150288: A Phase 2 Open-label Study Investigating the Safety and Efficacy of Binatumomab After Frontline R-Chemotherapy in Adult Subjects With Newly Diagnosed High-risk Diffuse Large B-Cell Lymphoma (DLBCL)	21/03/2017	Yes	25/01/2017	25/01/2017	25/01/2017	08/02/2017	13/02/2017	15/02/2017								
71	16/SC/0391	208610	A randomized, double-blind, double-dummy, parallel-group study comparing the efficacy and safety of ofatumumab versus teriflunomide in patients with relapsing multiple sclerosis.	12/04/2017	Yes	05/09/2016	24/02/2017	13/10/2016	24/02/2017	06/03/2017	15/03/2017								
72	17/NE/0115	218417	Multi-center, international, double-blind, two-arm, randomized, placebo-controlled phase II trial of pirfenidone in patients with unclassifiable progressive fibrosing ILD	31/07/2017	Yes	07/12/2016	01/06/2017	26/05/2017	14/06/2017	23/06/2017	23/06/2017								
73	17/WM/0216	225761	Investigating the benefits of a 3D camera for recording healing wound dimensions	27/07/2017	Yes	04/05/2017	29/06/2017	30/06/2017	10/07/2017	10/07/2017	10/07/2017								
74	16/EE/0463	214371	COMPLEMENT-1: An open-label, multicentre, Phase IIIb study to assess the safety and efficacy of ribociclib (LEE011) in combination with letrozole for the treatment of men and postmenopausal women with hormone receptor-positive (HR+) HER2-negative (HER2-) advanced breast cancer (aBC) with no prior hormonal therapy for advanced disease	26/04/2017	Yes	19/10/2016	20/02/2017	30/01/2017	22/02/2017	23/02/2017	28/03/2017								
75	16/YH/0506	212039	A Randomized, Double-Blind, Parallel Group, Multi-Centre Study to Assess the Efficacy and Safety of PT009 compared to PT005 on COPD Exacerbations over a 52-Week Treatment Period in Subjects With Moderate to Very Severe COPD	22/05/2017	Yes	10/01/2017	31/03/2017	22/02/2017	03/04/2017	11/04/2017	04/05/2017								
76	17/NS/0020	224422	Double blind randomised controlled trial of Remote Ischemic preconditioning in Multiple sclerosis	02/05/2017	Yes	28/12/2016	17/03/2017	17/03/2017	05/04/2017	05/04/2017	05/04/2017								
77	17/YH/0103	221203	Prolonged Enoxaparin in primary Percutaneous Coronary Intervention; a Pilot Pharmacodynamic study (PENNY PCI)	14/06/2017	Yes	13/03/2017	23/05/2017	23/05/2017	12/06/2017	12/06/2017	12/06/2017								

