

Research Ethics Committee Reference Number	Integrated Research Application System Number	Name of Trial	Date of Receipt of Valid Research Application	Date of First Patient Recruited	A - Permissions delayed/denied	B - Suspended by sponsor	C - Closed by sponsor	D - Sponsor Delays	E - Staff availability issues	F - No patients seen	G - No patients consented	H - Contracting delays	I - Rare diseases	J - Other	Comments	Reasons for delay correspond to:
16/LO/0019	180161	Bone Evaluation in women over 40 who Switch from Truvada/NNRTI to Triumeq	29/03/2016	11/07/2016	FALSE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	Low number of potential patients. <10% women in clinic cohort. Patients seen every 6 months to once a year.	Neither
13/EM/0073	120873	The Role of Glasses Wearing in Amblyopia Treatment. A randomised controlled multi-centre trial	29/03/2016		FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	PI has offered the study to several patients but they have not agreed to go on to the study. RN has made a recruitment plan which involves introducing the study at a different time point to ease burden for patients. No patients yet consented (as at 31.3.17).	Neither

15/EE/0421	191851	Pomalidomide in Relapsed and Refractory Multiple Myeloma	29/03/2016		FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	Site activation delayed by 21 days post-SIV due to Sponsor issues. Several patients screen failed/declined as at 31.03.17; subsequent change in treatment pathway means no longer as interesting for patients. Drug became available through the cancer drugs fund and so study became less attractive	Sponsor
16/EM/0007	190428	Evaluate Rivaroxaban in Patients after Successful Transcatheter Aortic Valve Implantation	29/03/2016	24/06/2016	TRUE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	SSI submitted early (29.3.16) due to changeover to HRA. Subsequently the sponsor did not issue the study drugs to pharmacy in time.	Sponsor
16/LO/0039	195230	A Phase 3, Randomized, Double-Blind Study to Evaluate the Safety and Efficacy of Switching from a Regimen containing Dolutegravir and ABC/3TC, or a Fixed Dose Combination (FDC) of ABC/DTG/3TC to a FDC of GS-9883/F/TAF in HIV 1 Infected Subjects who are Virologically Suppressed	29/03/2016	03/05/2016	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	70 day target met	Neither
14/LO/1559	152866	Sorin Universal Registry on Aortic Valve Replacement	29/03/2016	07/06/2016	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	70 day target met	Neither

16/LO/0240	199083	An open-label, prospective, non randomised, multicentre study to evaluate clear skin effect on health-related quality of life outcomes at 16 and 52 weeks in patients with moderate to severe plaque psoriasis treated with secukinumab 300 mg s.c. with or without previous exposure to systemic therapy	29/03/2016	03/05/2016	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	70 day target met	Neither
15/NW/0171	168748	A randomised, double blind, placebocontrolled trial of a twoweek course of dexamethasone for adult patients with a symptomatic chronic subdural haematoma	29/03/2016	27/05/2016	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	70 day target met	Neither
15/LO/0539	166304	A Randomised phase II trial of Adaptive Image guided standard or Dose Escalated tumour boost Radiotherapy in the treatment of transitional cell carcinoma of the bladder	29/03/2016	12/01/2017	TRUE	FALSE	FALSE	FALSE	TRUE	FALSE	TRUE	FALSE	FALSE	FALSE	SSI submitted early (29.3.16) due to changeover to HRA. Long delay in study activation due the QA requirement regarding radiographer training.	NHS Provider
15/LO/0928	170943	Mitral Valve Repair Clinical Trial	29/03/2016	26/07/2016	TRUE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	SSI submitted early (29.3.16) due to changeover to HRA. Subsequent to NHSP, the sponsor stated they needed to obtain MHRA approval, having previously said they didn't need it. This was issued mid-June but it wasn't possible to arrange an SIV until late July.	Sponsor

15/LO/2124	192795	A Randomized Multicenter Pivotal Study of CDX-011 (CR011-vcMMAE) in Patients with Metastatic, GPNMB Over-Expressing, Triple-Negative Breast Cancer	29/03/2016	20/09/2016	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	SSI submitted early (29.3.16) due to changeover to HRA. 6 patients consented for pre screening tissue analysis, 3 screen failed, 1 on study, 2 awaiting pre screen results to see if they are eligible.	Neither
GTAC182	64830	A Randomised Parallel Group Double-Blind Phase II Trial to Assess the Activity of TroVax® (MVA-5T4) Versus Placebo in Patients with Relapsed Asymptomatic Epithelial Ovarian, Fallopian Tube or Primary Peritoneal Cancer	29/03/2016	04/07/2016	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	SSI submitted early (29.3.16) due to changeover to HRA, SIV 31/5/16, study not activated until 21/6/16 due to missing delegation log information. H&S Executive approval also required for study set up. 1st patient recruited 4/7/2016.	NHS Provider
16/NI/0034	194752	The Medtronic CoreValve™ Evolut R™ FORWARD Study	29/03/2016	27/07/2016	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	SSI submitted early on 29.3.16 due to changeover to HRA Approval. We were not in a position to be able to issue R&D approval until 22/6/16. First patient recruited 26/7/16, so within target timeframe from NHSP.	Neither

15/EE/0435	191299	Stratified Treatment Optimisation for HCV-1 (STOPHCV-1)	29/03/2016	27/06/2016	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	SSI submitted early on 29.3.16 due to changeover to HRA Approval. We were not in a position to be able to issue R&D permission until 21/6/16, and first patient was subsequently recruited within 1 week.	Neither
15/LO/2087	145273	Prospective, Randomised, Fellow Eye Study evaluating correlation between Ciliary Sulcus Anatomy with other Ocular Parameters using Ultrasound Biomicroscopy after horizontal or vertical placement of the intraocular lens in the capsular bag during standard cataract surgery.	29/03/2016	12/07/2016	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	SSI submitted early on 29.3.16 due to changeover to HRA Approval. We were not in a position to be able to issue R&D permission until 16/6/16. First patient recruited 11/7/16, so within target timeframe from NHSP.	Neither
15/LO/0769	155035	Left Atrial Appendage Occlusion Study III	29/03/2016		TRUE	FALSE	FALSE	TRUE	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	Sponsor delay with setup, due to waiting for an amendment to go through. Delay initially at site as surgeon (PI) broke his hand. Practice has changed since we were approached, so we may not be able to recruit.	Both