

Research Ethics Committee Reference Number	Integrated Research Application System (IRAS) number	Name of Trial	Target number of patients agreed?	Min. number of patients agreed	Max. 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	Total number of study particip-ants recruited	Date that the trial closed to recruit-ment	Reason for closure of trial	Comments
14/SC/1161		Prospective, single-arm, multicentre, observational registry to further validate safety and efficacy of the ultimaster DES in real-world patients.	30	30		Not Available / Not Agreed	25/07/2017	0	21		Recruit- ment Finished	Global competition.
16/EM/0133		A randomized, double-blind, placebo-controlled multicenter study of secukinumab to evaluate the safety, tolerability and efficacy up to 2 years in patients with active nonradiographic axial spondyloarthritis	5	5		Date Agreed	01/09/2017	7	1		Recruit- ment	Three additional patients screen failed prior to randomisation. Lack of patients meeting inclusion criteria.
17/EM/0075		Evaluating the tolerance, compliance and acceptability of a nutritionally complete, high energy, high protein, enteral feed in adults ? a pilot study	5	5		Date Agreed			9		Recruit- ment Finished	Target met.

30/01/2019 1 of 3



Research Ethics Committee Reference Number	Integrated Research Application System (IRAS) number	Name of Trial	Target number of patients agreed?	Min. number of patients agreed	Max. 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	Total number of study particip-ants recruited	Date that the trial closed to recruit-ment	Reason for closure of trial	Comments
		A RANDOMIZED, DOUBLE-										
		BLIND, PLACEBO-CONTROLLED,										
		MULTICENTER LONG-TERM										
		SAFETY AND TOLERABILITY										
		STUDY OF ETC-1002 IN										
		PATIENTS WITH										
		HYPERLIPIDEMIA AT HIGH										
		CARDIOVASCULAR RISK WHO				Not						
		ARE NOT ADEQUATELY				Available					Recruit-	
		CONTROLLED BY THEIR LIPID-				/ Not					ment	
16/LO/0553	194625	MODIFYING THERAPY	8	8	8	Agreed			8	18/04/2018	Finished	Target met.
		A MULTICENTER OPEN-LABEL EXTENSION (OLE) STUDY TO ASSESS THE LONG-TERM SAFETY AND EFFICACY OF BEMPEDOIC				Not Available / Not	/00 /00 -				Recruit- ment	
17/YH/0071	223653	ACID (ETC-1002) 180 MG	3	3	3	Agreed	14/09/2017	6	3	23/03/2018	Finished	Target met.
		A randomized, double-blind, placebo-controlled, parallel-group, multicenter, event-driven phase 3 study to investigate the efficacy of finerenone on the progression of deterioration of kidney									Recruit-	High number of patients
		function in patients with type 2				Date					ment	failed screening tests post
15/SW/0194	185459	diabetes mellitus and the	12	12	12	Agreed	30/11/2018	2	8	08/06/2018	Finished	consent.

30/01/2019 2 of 3



Research Ethics Committee Reference Number	Integrated Research Application System (IRAS) number	Name of Trial	Target number of patients agreed?	Min. number of patients agreed	Max. 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	Total number of study particip-ants recruited	Date that the trial closed to recruit-ment	Reason for closure of trial	Comments
16/EM/0193		A phase III, double-blind, randomized placebo-controlled study to evaluate the effects of dalcetrapib on cardiovascular (CV) risk in a genetically defined population with a recent Acute Coronary Syndrome (ACS): The Dal-GenE trial	6	6	6	Date Agreed	24/03/2017	6	2			High number of screen failures.
15/SC/0448		A randomized, double-blind, placebo-controlled, parallel-group, multicenter, event-driven phase 3 study to investigate the efficacy of finerenone on the reduction of cardiovascular morbidity and mortality in patients with type 2 diabetes mellitus and	9	9	9	Date Agreed			10		Recruit- ment Finished	Target met.

30/01/2019 3 of 3