

Performance in Delivering studies Q4

Research Ethics Committee Reference Number	Integrated Research Application System Number	Name of Trial	Drop down options. Please indicate whether the trial has a fixed target (Number Agreed), range target (Range Agreed) or no target agreed or available (Not Available /Not Agreed)Target Number Of Patients Agreed?	Minimum Number Of Patients Agreed (Enter Same In Both If Only One Number)	Maximum Number Of Patients Agreed (Enter Same In Both If Only One Number)	Drop down options. Indicate if a target date has been agreed.Target Date To Recruit Patients Agreed?	Format dd/mm/yyyy. Please show the date agreed to recruit patients. Leave blank if unavailable.Date Agreed to recruit target number of patients	Please report the total number of patients recruited at the agreed target date.Total Number Of Patients Recruited At The Agreed Target Date	Please only include records for commercial trials which have closed to recruitment within the reporting window. Format dd/mm/yyyy.Date That The Trial Closed To Recruitment	Please report the total number of patients recruited.Total Number Of Study Participants Recruited	Reason For Closure Of Trial
13/EM/0459	137785	POSNOC - POSitive Sentinel NOde: adjuvant therapy alone versus adjuvant therapy plus Clearance or axillary radiotherapy. A randomised controlled trial of axillary treatment in women with early stage breast cancer who have metastases in one or two sen	Number Agreed	25	25	Not Available / Not Agreed			13/07/2021	28	Recruitment Finished- Sponsor reached their recruitment target.
16/SS/0014	187949	GEM3: A double blind placebo controlled trial of a combination of methotrexate and gefitinib versus methotrexate alone as a treatment for ectopic pregnancy	Number Agreed	1	1	Date Agreed	31/12/2019	5	08/10/2021	10	Recruitment Finished- End of study
19/SC/0333	254064	ObsQOR ObsQOR Study: Quality of Recovery from Obstetric Anaesthesia- a multicentre study.	Number Agreed	15	15	Date Agreed	22/10/2021	11	22/10/2021	11	Recruitment Finished- End of Study
18/WM/0132	237111	OPTI-SURF: Optimal surfactant delivery for preterm babies with respiratory distress syndrome. The effect of surfactant dose on outcomes in preterm infants with Respiratory Distress Syndrome.	Number Agreed	50	50	Not Available / Not Agreed			26/11/2021	67	The trial closed because it had reached the recruitment target.
19/SC/0295	256518	GWEP Study: A Randomized, Double blind, Placebo controlled Single ascending Dose Trial to Evaluate the Safety, Tolerability, and Pharmacokinetics of GWP42003-P in Conjunction with Hypothermia in Neonates with Moderate or Severe Hypoxic Ischemic Encep	Number Agreed	1	1	Not Available / Not Agreed			10/12/2021	0	Initially we managed to recruit and consent 3 babies that passed the pre-screening eligibility criteria, in compliance with the protocol. However, these babies then failed the post-screening test because their EEG test result wasn't within the post-screening inclusion criteria.