

Research Ethics Committee Reference Number	Integrated Research Application System (IRAS) number	Name of Trial	Date Site Invited	Date Site Selected	HRA Approval Date	Date Site Confirmed By Sponsor	Date Site Confirmed	Date Site Ready to Start	Date of First Patient Recruited	Non- Confirmation Status	Duration between Date site selected and First Patient (days)	Benchmark Met	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:
17/EM/0075	222172	Evaluating the tolerance, compliance and acceptability of a nutritionally complete, high energy, high protein, enteral feed in adults ? a pilot study		27/03/2017	27/03/2017	24/04/2017	02/05/2017	08/05/2017	17/05/2017		51	Yes						Y			Final complete costing template not received until 24/04/2017 from sponsor.	Both
16/HRA/5525	210175	MindSHINE 3: A definitive randomised controlled trial investigating two online wellbeing interventions to reduce NHS staff stress	17/01/2017	06/04/2017	23/01/2017	06/04/2017	08/05/2017	15/05/2017	06/06/2017		61	Yes										
17/YH/0071	223653	A MULTICENTER OPEN- LABEL EXTENSION (OLE) STUDY TO ASSESS THE LONG-TERM SAFETY AND EFFICACY OF BEMPEDOIC ACID (ETC-1002) 180 MG	03/04/2017	11/05/2017	19/07/2017	29/06/2017	04/07/2017	30/08/2017	01/09/2017		113	No	Y			Y	Y				HRA approval for the study was not received until 19/07/17 (initial HRA assessment letter dated 22/02/17) - cause of delay not known by site.2. Patients entering this extension study can only do so when they have fully completed the sister study. Our first eligible patient completing the sister study was seen and recruited to this study within the protocol required timelines.3. Sponsor could not provide an earlier date for the Site Initiation Visit.	
16/NW/0517	188554	A phase III study to determine the role of ixazomib as an Augmented Conditioning therapy in salvage autologous stem cell transplant (ASCT) and as a post-ASCT Consolidation and maintenance strategy in patients with Relapsed multiple myeloma.		05/04/2017	27/10/2016	08/08/2017	18/08/2017	18/10/2017				No				V	Y				Delay due to time taken for study review and consultation by local clinicians. Local information pack of documents was provided to site prior to site having a confirmed PI. Delay between DSC and Date Site Ready to Start was due to pharmacy staff capacity for electronic	,



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17/SC/0142	215503	Evaluating the clinical and cost- effectiveness of permissive hypotension in critically ill patients aged 65 years or over with vasodilatory hypotension	03/03/2017	11/05/2017	24/04/2017	18/08/2017	29/08/2017	01/09/2017	02/09/2017		114	No					Y					The sponsor required GCP training of staff at site prior to release of sponsor signed contract - staff availability issues to undertake GCP training at the NHS Provider caused a delay.2. Sponsor and NHS Provider could not find an early mutually agreeable date for the Site Initiation Visit which also created a delay.	
16/NE/0400	212212	A Randomised Controlled Trial of the effect of a Two-layer Compression Bandage System on Knee Function following total knee arthroplasty	19/05/2017	28/09/2017	08/02/2017	28/09/2017	07/11/2017	09/11/2017	16/11/2017		49	Yes											
17/LO/0365	216598	Improving Fundamental Care on Hospital Wards	28/07/2017	28/07/2017	28/04/2017	23/08/2017	08/09/2017	24/10/2017	06/11/2017		101	No				Y						Overall delays due to sponsor staff annual leave. DSS to DSC delay due to sponsor not sending contract through to NHS Provider in a timely manner. DSC to FPR delay due to sponsor not submitting to HRA for protocol amendment in a timely manner and requirement to wait for approval.	Sponsor

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