Research			Reasons for not achieving the 70 day target from receipt of valid research application to 1st patient recruited											Reasons for	
Ethics Committee Reference 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	delay correspond to:
13/LO/0908	A Phase 2, Single-Arm, Open-Label, Multicenter Study of the Clinical Activity and Safety of Enzalutamide in Patients With Advanced, Androgen Receptor- Positive, Triple-Negative Breast Cancer	16/01/2014	19/03/2014											70 day target met	
12/EE/0445	A randomised double-blind controlled phase III study to compare the efficacy and safety of intravenous ferric carboxymaltose with placebo in patients with anaemia undergoing major open abdominal surgery	20/01/2014	05/06/2014							Y				Every effort is going into recruiting to this study, but recruitment is difficult due to the haemoglobin range required, and the short notice given pre-surgery (c.10 days). Research bloods are required at least 10 days before surgery, which means there is often no time for consent. Other Trusts are experiencing the same problem with this study.	Neither
13/SC/0016	Randomized open label study of oral versus intravenous antibiotic treatment for bone and joint infections requiring prolonged antibiotic treatment: Multi- centre study	06/02/2014	08/04/2014											70 day target met	
11/WS/0118	Safety and Efficacy assessment of Monoprost® (unpreserved latanoprost) in comparison with Lumigan® 0.01 % and Lumigan® 0.03% UD, in patients with primary open angle glaucoma or ocular hypertension, stabilized by Lumigan® 0.01 % with ocular surface intoler	14/02/2014	23/04/2014											70 day target met	
13/EW/0349	A Randomized Phase II Study of Fulvestrant in Combination with the dual mTOR Inhibitor AZD2014 or Everolimus or Fulvestrant alone in Estrogen Receptor- Positive Advanced or Metastatic Breast Cancer.	10/02/2014	23/10/2014	Y	Y					Y				This study is only expected to recruit 3 patients over 2 years, which equates to 1 patient every 8 months. It is therefore not surprising that there were no patients recruited within the FPR target timeframe. Every effort is being made to recruit, and the 1st patient was recruited 23/10/14. 3 further patients failed screening; 1 further patient was consented but subsequently failed eligibility due to a liver reading.	Neither
13/YH/0389	52 WEEK, PHASE 3 DOUBLE-BLIND, RANDOMIZED, PLACEBOCONTROLLED, PARALLEL- GROUP STUDY TO ASSESS THE EFFICACY, SAFETY AND TOLERABILITY OF PF-04950615 IN SUBJECTS WITH HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA.	19/03/2014	. N/A				Y			Y				Sponsor delayed site activation. Not activated until 56 days after the SIV, just 4 working days before FPR target date. As at 29/07/2014, 137 sets of patient notes had been screened for eligibility; some patients had de	Sponsor
13/LO/1795	A double-blind, randomized, placebo- controlled, cross-over study to evaluate the clinical efficacy and safety of subcutaneous administration of human plasma-derived C1-esterase inhibitor in the prophylactic treatment of hereditary angioedema		02/06/2014							Y				Three patients were approached within target period, but they declined to participate. BSUH was the first site in Europe to randomise a patient.	Neither
12/LO/1078	A phase II randomised trial of carfilzomib, cyclophosphamide and dexamethasone (CCD) vs cyclophosphamide, velcade and dexamethasone (CVD) for first relapse and primary refractory multiple myeloma	25/03/2014	29/05/2014											70 day target met, despite being unable to approach patients until end of May due to IP not being available.	

	1	1	Ī	1	1					1	1
14/EM/0032	A 52-week, multicenter, randomized, double-blind study of subcutaneoussecukinumab to demonstrate efficacy as assessed by Psoriasis Area and Severity Index at 16 weeks of treatment compared to ustekinumab and to assess long-term safely, tolerability and efficacy in subjects with moderate to severe plaque psoriasis	02/05/2014	N/A		Υ					This study was given NHSP on the same day as VRA but was closed by the Sponsor 32 days later as their global recruitment target was met. Every effort was made to recruit, but we were unable to recruit within this short timescale.	Neither
14/LO/0440	Prospective Randomised Controlled Trial comparing Monofocal Intraocular Lenses and Limbal Relaxing Incisions with Toric Intraocular Lenses for correcting Astigmatism up to 2.5 Diopters during standard cataract surgery.	03/06/2014	11/06/2014							70 day target met	
13/LO/1720	A Phase 2, Randomized, Double Blind, Placebo Controlled, Multicenter Study of Efficacy and Safety of Enzalutamide in Combination With Exemestane in Patients With Advanced Breast Cancer That Is Estrogen or Progesterone Receptor Positive and HER2 Normal	05/06/2014						Υ		This is a difficult study to recruit to, and as at December 2014 we have had 3 screen failures and 1 patient who declined to participate.	Neither
14/LO/0083	An open label study examining the efficacy and cardiovascular risk of immediate versus deferred switch from a boosted P1 to dolutegravir (DTG) in HIV infected patients with stable virological suppression. NEAT 22 /SSAT 060	02/05/2014	04/07/2014							70 day target met	
14/SC/0225	Gilead 311-1089 Phase 3 randomised open-label switch study to evaluate F/TAF in HIV-1 positive subjects who are virologically suppressed on regimens containing FTC/TAF (GILEAD 1089)	16/06/2014	22/07/2014							70 day target met	
13/NS/0143	The SIMS trial	13/06/2014	04/09/2014					Y		Every effort was made to achieve the FPR target, with screening at the fortnightly clinic run by the PI starting as soon as the study was approved. Many patients did not consent or were not eligible (NB 8 patients were screened and ineligible before the first patient was recruited). There are only 3 or 4 patients per clinic, and no clinics have been missed by the research nurse team.	Neither
13/LO/1426	An international multicentre open-label comparative therapeutic exploratory trial to investigate the role of a new neonatal formulation of dobutamine in the treatment of haemodynamic insufficiency in the immediate postnatal period.					Υ				The Sponsor delayed opening BSUH as a site. We were only activated on 1/8/14, 37 days past the FPR target date.	Sponsor
13/YH/0282	ACT-MOVE: ML28641 - Subcutaneous tocilizumab in rheumatoid arthritis	16/04/2014 31/03/2014					Y			Every effort is being made to recruit to this study, but it is a difficult trial to recruit to as the study drug was licensed shortly afer the study opened, and so is available on prescription without the need to be a research participant.	Neither
13/EE/0126	Evaluation of Safety and Efficacy of the BACE™ [Basal Annuloplasty of the Cardia Externally] Device in the Treatment of Functional Mitral Valve Regurgitation [FMR]  A Phase IIIb, Multi-Center, Randomized,	04/06/2014	-						Y	No patients recruited by target date, which was expected as this is a rare disease category study.	Neither
14/LO/0081	Double-Blind, Placebo-Controlled, Multidose, 24-Week Study to Evaluate the Efficicacy and Safety of Atacicept in Subjects With Systemic Lupus Erythematosus	23/05/2014	<u>-</u>					Υ		Difficult study to recruit to. As at December 2014 we have had one screen failure and one patient declined.	Neither
13/LO/1082	Revascularisation or medical therapy in elderly patients with acute anginal syndromes.	06/05/2014	13/06/2014							70 day target met	

	Does oral sodium bicarbonate therapy improve function and quality of life in													
	older patients with chronic kidney disease												70 day target met	
12/ES/0023	and low-grade acidosis?	23/06/2014	21/08/2014											
	A Prospective, Single-Arm, Clinical- Setting Study to Describe Efficacy,												70 day target met. BSUH has over-recruited to this study, and	
	Tolerability and Convenience of												is the top recruiting site within	
	Teriflunomide Treatment Using Patient												Kent, Surrey and Sussex for	
14/NW/0017	Reported Outcomes (PROs) in Relapsing Multiple Sclerosis (RMS) Patients.	22/04/2014	10/06/2014										dementia trials.	
14/14///0017	Multiple deletosis (NMO) Fatterits.	22/04/2014	10/00/2014										PI is screening but has not	
	REstart or STop Antithrombotics							Y					been able to identify any	
12/SS/0138	Randomised Trial (RESTART).  Optimisation and Individualisation of	14/03/2014	-										eligible patients	Neither
	HeartSparing Breast Radiotherapy													
	Techniques (The HeartSpare Study):												70 day target met	
13/LO/0181	Stage II	27/03/2014	02/06/2014											
	PROSPER: A Multinational, Phase 3,												BSUH was unable to be opened as a site until 1/9/14	
	Randomized, Double-Blind, Placebo-												due to unsigned paperwork	
	Controlled, Efficacy and Safety Study of			Y					Y				required by Sponsor. As at 31	
	Enzalutamide in Patients With Nonmetastatic Castration-Resistant												December 2014 there have been 5 screen failures and 2	
13/LO/1081	Prostate Cancer	18/07/2014	N/A										patients have declined.	NHS Provider
10/20/1001		10/07/2011											patiente nave decimica.	TWIGHT TOVIGO
	SAPROCAN: Saracatinib (AZD0530) and													
	docetaxel in metastatic, castraterefractory prostate cancer: a phase I/randomised												70 day target met	
	phase II study by the UK													
11/AL/0081	NCRI Prostate Clinical Studies Group	18/07/2014	05/08/2014											
	An exploratory, randomised, double-blind, controlled study to assess the effect of an													
	Amino Acid Based Formula with a													
	synbiotic blend on gut microbiota and												70 day target met	
	stool characteristics in infants with													
13/NE/0126	suspected gastrointestinal Non IgE mediated cow's milk allergy (CMA).	20/06/2014	14/08/2014											
10/112/0120	mediated cow o mink dilongy (com t).	20/00/2011	11/00/2011											
	A Prospective, Randomized Evaluation of													
	the TriGuard™ HDH Embolic DEFLECTion Device during Transcatheter												70 day target met.	
14/SW/0079	Aortic Valve Implantation	09/07/2014	17/07/2014											
	·													
	The United Kingdom Transcatheter Aortic													
	Valve Implantation (UK TAVI) Trial. A multi-centre randomised controlled trial to												BSUH only had clearance to start recruiting on 8/10/14,	
	assess the clinical effectiveness and cost-					.,						.,	which was after the FPR target	
	utility of TAVI, compared with					Υ						Υ	date. Delay caused by initial	
	conventional surgical aortic valve replacement, in patients with severe												'test' echo being rejected.	
	symptomatic aortic stenosis at													
13/LO/0451	intermediate or high operative risk.	01/07/2014	17/10/2014											Sponsor
	ABSORB UK Registry - A post-market													
	registry of patients with de novo lesions in previously untreated vessels treated with												70 day target met	
13/EM/0476	Absorb BVS	12/05/2014	29/05/2014											
													Laboratory manual required	
													clarification. Delays in response from Evelina Childrens Hospital	
	The Hypertension Optimal Treatment in Children with Chronic Kidney Disease												regarding use of their	
	study: The HOT-KIDS study- A					Υ			Y			Y	equipment at BSUH. As at the	
	randomised trial to compare effects of												end of December 2014,	
	aggressive versus standard targets in blood pressure on target organ damage in												multiple screen failures and 1 eligible patient declined study	
10/H0802/13	children with CKD.	31/07/2014	N/A		I				I	]		I	entry.	Sponsor
													Every effort is being made to	
					I				I	]		I	recruit, but no suitable patients	l
					1				1			1	have been found. Patients that	
					1				Υ			1	were initially eligible, subsequently failed screening	
	A phase III trial comparing standard				I				I	]		I	as they had co-morbidities that	
	versus novel CRT as preoperative treatment for MRI defined locally				I				I	]		I	precluded randomising them to	
10/H0706/65	advanced rectal cancer.	14/08/2014	N/A		1				1			1	potentially receive Irinotecan.	Neither
		. 1/00/2014			1	l	l		1		l	1		
	A Randomised Multicentre Accelerated				1				1			1		
	Radiotherapy Study of Dose Escalated Intensity Modulated Radiotherapy vs				I				I	]		I		l
	Standard Dose Intensity Modulated				I				I	]		I	70 day target met	l
	Radiotherapy in Patients Receiving				I				I	]		I		
10/H0715/48	Treatment for Locally Advanced Laryngeal and Hypopharyngeal Cancers.	14/08/2014	24/00/2044		1				1			1		
		14/00/2014	27/03/2014		1	1	1	1	1	I	1	1	1	

	RANDOMISED, DOUBLE-BLIND, PLACEBO-CONTROLLED (PART A) AND DOUBLE-BLIND, DOUBLE-DUMMY, ACTIVE-CONTROLLED (PART B),					Y			Y			needed to carry out all the screening assessments. Final training was given on 6/10/14, only 9 days before the FPR	Spons
	PARALLEL GROUP STUDY TO EVALUATE THE EFFICACY AND SAFETY OF RPC1063 ADMINISTERED		13/01/2015			ľ			1			target date. As at 31 December 2014 there had been 15 screen	'
	ORALLY TO RELAPSING MULTIPLE		(don't submit in Jan 2015									fails and one patient declined to participate.	)
4/EM/0129	SCLEROSIS PATIENTS	06/08/2014	report)										
	Open-Label Extension Study of EFC12492, R727-CL-1112, EFC12732, & LTS11717 Studies to Assess the Long-											70 day target met	
	Term Safety and Efficacy of Alirocumab in Patients with Heterozygous Familial												
4/WS/0004	Hypercholesterolmia A STUDY TO DETERMINE THE	12/08/2014	30/09/2014										
4/EE/1016	ACCURACY OF ZERO-FLUX AND INGESTIBLE THERMOMETERS IN THE PERIOPERATIVE SETTING	22/08/2014	24/09/2014									70 day target met	
	A Phase 1 Study to Evaluate the Safety and Tolerability of MEDI4736 in												
	Subjects with Myelodysplastic Syndrome after Treatment with								Y			One screen failure (5/11/14)	Neith
4/LO/0892	Hypomethylating Agents A Phase II, Double Blind, Randomized,	17/09/2014	N/A										1
	Placebo-Controlled Study of the AKT Inhibitor AZD5363 in Combination With Paclitaxel in Triple-Negative Advanced or											70 day target met	
4/LO/0121	Metastatic Breast Cancer COgnitive behavioural therapy vs	22/09/2014	29/09/2014										
	standardised medical care for adults with Dissociative non-Epileptic Seizures: A											70 day target met	
13/LO/1595	multi-centre randomised controlled trial (CODES).	17/10/2014	12/11/2014										
0	Reformulated raltegravir q.d. (1200 mg) versus raltegravir b.i.d. (400 mg) in ART-	00/40/0044	40/44/9044									70 day target met. First patient screened 13/11/2014. Study closed to recruitment and fully	
4/LO/1381	naïve pts (ONCE)	22/10/2014	13/11/2014									recruited target of 5 As at 31/12/14 - 10 screened. 2	
4/ES/0072	How can we optimise inhaled beta2 agonist dose as 'reliever' medicine for wheezy pre-school children.	15/10/2014										were eligible but 1 declined, and 1 got chickenpox so currently unable to participate in study.	
4/13/00/2	wheezy pre-school children.	13/10/2014										Four screen failures as at	
3/LO/0699	Evaluating the effects of novel GLP-1 analogue, liraglutide, in patients with Alziemer's disease (ELAD study)	30/10/2014							Y			December 2014. 2 potential screens need PI approval. Still within FPR target timeframe.	
	A randomized, Open Label, Multi-Centre,												
	Controlled Study to Assess Safety and efficacy of ELAD in subjects with Acute											No patients recruited yet. 70	
3/YH/0147	Alcoholic Hepatitis (AAH) Who have failed Steroid Therapy (incorporating VTI-210E as a follow up registry)	12/11/2014										day target is 21.01.2015	
	A Randomized, Open Label, Phase 4												
	Study Evaluating the Renal Effect of Elvitegravir/Cobicistat/Emtricitabine/Tenof												
	ovir DF or other Tenofovir DF-containing Regimens (Ritonavir-boosted Atazanavir											(Gilead) delayed study activation due to delay with	
	plus Emtricitabine/Tenofovir DF or Efavirenz /Emtricitabine/Tenofovir DF)											treatment arm drug. Still within 70-day target timeframe.	
	compared to Ritonavir boosted Atazanavir plus Abacavir/Lamivudine in Antiretroviral												
4/LO/1513	Treatment-naïve HIV-1 Infected Adults with eGFR ≥70 mL/min	23/10/2014				Y			١ ,				
720/1010	Mar oor it are including	20/10/2011										Site activation delayed post-SIV	,
												as no contract was in place.  Once this was resolved, the	
												Sponsor issued an amendment,	,
												so recruitment could not commence until the	
	A Phase III Trial of Surgery versus Active											amendment was processed and given all relevant approvals.	1
4/WM/0083	Monitoring for Low Risk Ductal Carcinoma in Situ (DCIS) (LORIS)	01/10/2014		Υ		Y			Y	Y			
			1 ]	Ī			1			1		Every effort has been made to recruit to this study, but 4	
	RIAItO: A Randomised Investigation of							1	1		İ	patients have failed screening	1

	A Prospective Randomised Phase III														
	Study of Observation Versus Screening MRI And													Many patients have been	
	Pre-Emptive													screened but none have been	
	Treatment in Castrate Resistant Prostate													eligible. Screening method	
	Cancer													being reviewed.	
12/LO/1109	Patients With Spinal Metastasis	17/10/2014								Y	·				Neither
														Still within FPR target	
	FOCUS4 - Molecular selection of therapy													timeframe. Not yet open to recruitment due to delays with	
	in colorectal cancer: a molecularly													study documentation.	
	stratified randomised controlled trials													study documentation.	
13/SC/0111	programme	24/10/2014		Y			Y								Both
	A multicenter, Single Arm Study of														
	Enzalutamide in Patients with Progressive														
	Metastatic Castration-Resistant Prostate													70 day target met.	
14/LO/0298	Cancer Previously Treated With Abiraterone Acetate.	30/10/2014	25/11/2014												
14/LO/0298	Abiraterone Acetate.	30/10/2014	25/11/2014												
	A Phase 2, Randomized, Open-Label,				1	1	1	1		1		1	1		
	Parallel Group Study Evaluating the														
	Safety andEfficacy of TAK-385, an Oral														
	Gonadotropin-Releasing Hormone													No patients recruited yet. 70	
	(GnRH) Antagonist, forPatients With													day target is 27.01.15	
	Localized Prostate Cancer Requiring														
	Neoadjuvant and Adjuvant														
	AndrogenDeprivation Therapy With														
14/LO/1052	External Beam Radiation Therapy (EBRT)	18/11/2014													
	A Phase III, Randomized, Double-Blind,														
	Placebo-Controlled, Multicenter Clinical														
	Study Evaluating the Safety and Efficacy													No patients recruited yet. 70	
	of Icatibant as a Treatment for													day target is 02.03.15	
	Angiotensin-Converting Enzyme Inhibitor														
14/EM/1070	(ACE-I)-Induced Angioedema in Adults	22/12/2014													
	Phase I study of KHK2823 in Patients													No patients recruited yet. 70	
	with Acute Myeloid Leukaemia or													day target is 25.02.15.	
14/YH/008	Myelodysplastic Syndrome Multi-centre Randomised Controlled Trial	17/12/2014												,g	
	of Angiotensin Converting Enzyme														
	inhibitor (ACEi)/Angiotensin Receptor													70 day target met	
	Blocker (ARB) withdrawal in adnoace													70 day target met	
13/YH/0394	renal disease: The STOP-ACEi Trial	14/10/2014	14/11/2014												
	RESPOND: Repositionable Lotus Valve				İ		İ	İ	İ	İ		İ			
	System-Post Market Evaluation of Real													70 day target met	
14/YH/0086	World Clinical Outcomes	06/10/2014	22/10/2014											1	
	Prospective, single-arm, Multi-centre, observational registry to Further Validate														
	Safety and Efficacy of the Ultimaster DES													70 day target met	
	system in unselected patients														
14/SC/1161	representing everyday clinical practice	18/12/2014	22/12/2014												
	, , , , , , , , , , , , , , , , , , , ,				İ		İ	İ	İ	İ		İ			
1					ĺ		ĺ	ĺ	ĺ	ĺ		ĺ			
1	A Randomized, Double blind, Placebo-														
	Controlled, 2-Part Study of Orally													No patients recruited yet; IMP	
	Administered ALS-008176 to Evaluate the													still not received at site as at 31	
	Safety, Tolerability, Pharmacokinetics and													December 2014. 70 day target	
1	Pharmacodynamics of Single Ascending Dosing and Multiple Ascending Dosing in													is 17.02.15.	
1	Infants Hospitalized with Respiratory														
14/WM/0013	Syncytial Virus (RSV) Infection.	19/12/2014					·			· ·					Sponsor
		10,12,2014	07/01/2015				· '			<u> </u>				o.:. :	Оролоот
	More Response on Cardiac		(to include in											Still within FPR target timeframe as at 31 December	
	Resynchronization Therapy (CRT) with		next											timerrame as at 31 December 2014.	
13/LO/1891	MultiPoint Pacing (MPP)	04/11/2014	submission)			1					1			2017.	