Research Ethics Committee Reference Number	IRAS number	Full Name of Trial	Target number of patients available?	Minimum number of patients agreed to be recruited	Maximum number of patients agreed to be recruited	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 patients recruited when recruitment ended	Date trial closed to recruitment	Reason for the closure of the trial
11/LO/0537	75448	A Multicenter, Randomized, Double-Blind, Placebo- Controlled Study of the Efficacy of Natalizumab on Reducing Disability Progression in Subjects With Secondary Progressive Multiple Sclerosis	Number agreed	6	6	Date agreed	01/02/2013	5	5	24/11/2015	Recruitment finished
, , ,		Phase 4 trial comparing cumulative incidence of SCC after treatment + ingenol mebutate & imiquimod for multiple actinic keratoses on face &scalp. A multi-centre, randomised, two-arm, open label, active-controlled, parallel									
13/NW/0316	124757	group, 36-month trial. Reformulated raltegravir q.d. (1200 mg) versus raltegravir	Number agreed	8	8	Date agreed	31/05/2015	10	10	31/03/2016	Recruitment finished
14/LO/1381	153109	b.i.d. (400 mg) in ART-naïve pts	Number agreed	5	5	Date agreed	30/04/2015	5	5	05/12/2015	Withdrawn by Sponsor
14/SS/1048		A Phase 3b, Multi-center, Randomized-withdrawal, Placebo- controlled, Double blind, Parallel-group Trial to Compare the Efficacy and Safety of Tolvaptan (45 to 120 mg/day, Split-dose) in Subjects with Chronic Kidney Disease Between Late Stage 2 to Early Stage 4 Due to Autosomal Dominant Polycystic Kidney Disease	Number agreed	5	5	Date agreed	10/07/2015	5	5	01/03/2016	Recruitment finished
		A Phase 3, randomized, active-controlled, open-label study to evaluate the efficacy, safety and tolerability of switching to a darunavir/cobicistat/emtricitabine/tenofovir alafenamide (D/C/F/TAF) noce-daily single-tablet regimen versus continuing the current regimen consisting of a boosted protease inhibitor (bPI) combined with emtricitabine/tenofovir disoproxil fumarate (FFC/TDF) in virologically-suppressed, human immunodeficiency disoproxil fumarate (FFC/TDF) in virologically-suppressed, human immunodeficiency turns type 1 (HIV-1) infected									
14/WM/1210	164208	subjects.	Number agreed	4	4	Date agreed	12/08/2015	11	11	29/01/2016	Recruitment finished
14/NE/1185	152820	Short duration of dual antiplatElet therapy with SyNergy* II everolimus eluting stent in patients Older than 75 years undergoing percutaneous coronary Revascularization. The SENIOR trial	Range agreed	15	40	Date agreed	30/06/2015	7	7	11/02/2016	Recruitment finished
13/EM/0348	122371	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 coular hypertension, stabilized by Lumigan® 0.01 % with ocular surface intolerance	Number agreed	6	6	Date agreed	30/04/2016	2	2	30/04/2016	Recruitment finished
		Comparative Testing of 3 mL TransFix/EDTA Vacumm Blood Collection Tubes (TVTs) and Cyto-Chex 5 mL Blood									
14/NW/1531	165328	Collection Tubes (BCTs) Part 1: Equivalence Study	Number agreed	10	10	Date agreed	05/11/2015	13	13	19/04/2016	Recruitment finished
15/YH/0045	168195	ABLATOR Ablation Observational Registry	Range agreed	25	100	Date agreed	17/07/2017	36	36	28/04/2016	Recruitment finished
		COOL-AMI EU CASE SERIES CLINICAL STUDY: a single-centre case series clinical study to assess the feasibility of integrating therapeutic hypothermia(TH)using the ZOLL IVTM System as an adjuvant therapy in percutaneous coronary intervention (PCI) in patients with acute myocardial infarction (AMI)									Withdrawn by Sponsor
13/EE/0335	13/965	Randomised Evaluation of dabigatran etexilate Compared	Range agreed	5	,	Date agreed	28/02/2016	4	4	20/02/2016	withurawn by Sponsor
15/SC/0280	170452	to warfarIn in pulmonaRy vein ablation: assessment of an uninterrupted periproCedUral antIcoagulation sTrategy (The RE-CIRCUIT Trial)	Range agreed	9	20	Date agreed	31/08/2016	8	8	29/06/2016	Recruitment finished

	A Phase 3, randomized, active-controlled, double-blind study to evaluate efficacy and safety of darunavir/cobicistat/emtricitabine/tenofovir alafenamide (D/C/F/TAF) once daily fixed dose combination regimen versus a regimen consisting of darunavir/cobicistat fixed dose combination coadministered with							
	emtricitabine/tenofovir disoproxil fumarate fixed dose							
15/LO/1239	combination in antiretroviral treatment-naïve human 184169 immunodeficiency virus type 1 infected subjects.	Number agreed		5 Date agreed	31/03/2016		02/02/2016	Withdrawn by Sponsor
13/10/1235	184109 Illimunouenciency virus type 1 illiecteu subjects.	Number agreed	3	5 Date agreed	31/03/2010	3	02/02/2010	Withurawn by Sporisor
15/EE/0322	A phase 2, double-blind, randomized, placebo-controlled study to investigate possible drug-drug interactions 183313 between clobazam and cannabidiol (GWP42003-P)	Number agreed	3	3 Date agreed	29/02/2016	1	10/06/2016	Recruitment finished
	, , , , , , , , , , , , , , , , , , , ,						,,,,	
16/LO/0036	A Phase 3, Randomized, Double-Blind Study to Evaluate the Safety and Efficacy of 65-9883/Emtricitabine/Tenofovir Alafenamide Versus Abacavir/Dolutegravir/Lamivudine in 195359 HIV-1 Infected, Antiretroviral Treatment-Naïve Adults	Necks		6 Not available / not agreed			40/05/2016	Withdrawn by Sponsor
16/LU/0036	195359 HIV-1 Infected, Antiretroviral Treatment-Naive Adults	Number agreed	ь	6 Not available / not agreed		5	10/06/2016	withdrawn by Sponsor
	A Phase 3, Randomized, Open Label Study to Evaluate the Safety and Efficacy of Switching from Regimens Consisting of Boosted Atazanavir or Darunavir plus either Emtricitabine/Tenofovir or Abacavir/Lamiwudine to GS- 9883/Emtricitabine/Tenofovir Alafenamide in Virologically							
16/LO/0026	195795 Suppressed HIV-1 Infected Adults	Number agreed	4	4 Not available / not agreed	+	3 3	22/06/2016	Withdrawn by Sponsor
	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 VI Infected Subjects who are Virologically							
16/LO/0039	195230 Suppressed	Number agreed	5	5 Not available / not agreed		2	03/06/2016	Withdrawn by Sponsor
			50		2.1.010015			
13/NE/0336	137605 Trial of Proton Pump Inhibitors in Throat Symptoms A Multicentre, Single-Blind, Randomised Parallel-Group Study to Assess the Short and Long-Term Efficacy of Certolizumab Pegol plus Methotrexate Compared with Adalimumab plus Methotrexate in Subjects with Moderate to Severe Rheumatoid Arthritis Responding Inadequately to	Number agreed	30	00 Date agreed	01/10/2017	9	12/02/2016	
12/NE/0198	100141 Methotrexate	Range agreed	3	5 Not available / not agreed		1 1	13/01/2016	Withdrawn by Sponsor
13/SC/0279	A Phase 3, Open-Label Study to Evaluate Switching from a TDF-Containing Combination Regimen to a TAF-Containing Combination Single Tablet Regimen (STR) in Virologically 124704 Suppressed, HIV1 Positive Subjects.	Number agreed	4	4 Not available / not agreed		4	16/06/2016	Recruitment finished
2,23,32,3	A Phase 3, Randomized, Open-Label, Active-Controlled			, aramana, not agreed			10,00,2010	
14/LO/1443	Study to Evaluate the Efficacy and Safety of FG-4592 in the Treatment of Anemia in End Stage Renal Disease Subjects on Stable Dialysis Converted from Epoetin or 149317 Darbepoetin Alfa Treatment	Number agreed	4	4 Date agreed	19/12/2017	1	28/04/2016	Recruitment finished
15/LO/0684	Effects of ODM-109 on respiratory function in patients with ALS. A randomised, double blind, placebo-controlled, crossover, 3-period, multicentre study with open-label follow-up							
	177109 extension	Number agreed	5	5 Date agreed	13/10/2015	4	27/07/2016	Recruitment finished
15/LO/0881	A Phase III Multicenter, Double Blind, Randomized, Active Comparator-Controlled Clinical Trial to Evluate the Safety and Efficacy of MK-149A One Daily Versus ATRIPLA Once- 177217 Daily in Treatment Naïve HIV - 1 Infected Subjects	Number agreed	5	5 Date agreed	28/02/2018	5	08/08/2016	Recruitment finished
13/10/0001	277227 Daily in Treatment Naive Filv - 1 infected Subjects	ramoer agreed		J J J J J J J J J J J J J J J J J J J	20/02/2010		00/00/2010	nea dieniene milistied
15/LO/1324	181497 AMPLATZER Amulet Observational Post-Market Study	Number agreed	10	.0 Date agreed	17/11/2019	19 19	24/08/2016	Recruitment finished
-0/20/2027	privin a meen randict addervational Lost-Market Study	scr agreea	101		11/11/2017	101	24,00,2010	diemene milaneu

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	ARMOR3-SV: A Phase 3, Randomized, Open-Label, Multi-								
	Center, Controlled Study of Galeterone Compared to Enzalutamide in Men Expressing Androgen Receptor Splice								
	Variant-7 mRNA (AR-V7) with Metastatic (M1) Castrate								
15/LO/0564	173148 Resistant Prostate Cancer (CRPC)	Number agreed	2	2 Date agreed	01/02/2017	0	0	26/07/2016 Withdra	awn by Sponsor
	Randomised phase 3 trial of enzalutamide in first line								
14/LO/2218	androgen deprivation therapy for metastatic prostate 161762 cancer.	Number agreed	4	4 Date agreed	31/03/2016	0	0	25/07/2016 Withdra	awn by Host
	Surveying People Experiencing Young Adult Kidney Failure								
15/SW/0101	159847 (SPEAK)	Number agreed	10	10 Not available / not agreed		9	9	30/09/2016 Recruitr	ment finished
14/WM/1128	150126 Prognostic biomarkers in HCV cirrhosis	Number agreed	20	20 Date agreed	30/12/2016	15	15	01/07/2016 Withdra	h C
14/ WW/1128	130126 Prognostic Diomarkers in new cirriosis	Number agreed	20	to Date agreed	30/12/2016	15	15	01/07/2016 Withdra	awii by Spoilsor
	Investigation of the clinical, serological and genetic factors that determine primary non-response, loss of response and								
	adverse drug reactions to Anti-TNF drugs in patients with								
12/SW/0323	115956 active luminal Crohn's Disease.	Number agreed	13	13 Date agreed	31/12/2016	18	18	06/07/2016 Recruitr	ment finished
	[
11/NW/0548	A Randomised Investigation of Alternative Ofatumumab- 76092 containing regimens in less fit patients with CLL	Number agreed	5	5 Date agreed	30/11/2015	2	2	18/07/2016 Withdra	awn by Host
	Plasma Exchange and glucocorticoid in anti_neutrophil	agreed		-1	30/11/2013			20/0//2020 Withdia	= 1 11031
00/110700/50	cytoplasm antibody associated vasculitis: a randomised 27957 controlled trial	Number	2	2 Data served	01/00/2016		_	20/00/2016	anna finish ad
09/HO709/56	2/95/ controlled trial	Number agreed	5]	3 Date agreed	01/09/2016	5	5	30/09/2016 Recruitr	ment finished