

Performance in Delivering Clinical Research - Quarter 2 (2014/15)

All hosted, commercial clinical trials active between 01 October 2013 - 30 September 2014

| Trust Reference Code | Research Ethics Committee Reference Number | Name of Trial | Recruitment Target | Date Agreed to Recruit Target Number of Patients | Trial Status | Target Met Within Agreed Timeframe? | Comments |
|----------------------|--|---|--------------------|--|-----------------------|-------------------------------------|---|
| C&W11/075 | 11/SC/0329 | A Phase 3, Randomised, Double-blind, Placebo-controlled, Parallel-group, Multi-centre Study to Evaluate the Safety and Efficacy of Ustekinumab Maintenance Therapy in Subjects with Moderately to Severely Active Crohn's Disease | 5 | No date agreed with sponsor | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W11/100 | 11/LO/1034 | A randomised, prospective study, assessing changes in cerebral function in treatment naive HIV-1 infected subjects commencing either boosted atazanavir with Truvada or boosted dolutravir with maraviroc and Kivexa | 7 | No date agreed with sponsor | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W12/092 | 12/LO/1434 | Lung Volume Reduction Coil Treatment in Patients with Emphysema (RENEW) Study | 8 | 30/10/2014 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W13/015 | 11/LO/1455 | Gilead HCV Registry 0122 Responders | 8 | 01/10/2016 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W13/068 | 13/EE/0241 | Secukinumab In patients with moderate to severe active, chronic plaque psoriasis who have failed on TNFa antiGoNists: A clinical Trial Evaluating Treatment Results (SIGNATURE) | 2 | 16/01/2015 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. Study recruitment window nationally extended from 09/10/2014 to 16/01/2015. |
| C&W13/085 | 13/EE/0270 | A Global Registry to Evaluate Long-Term Effectiveness of Neurostimulation Therapy for Pain | 30 | 04/07/2016 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W12/074 | 12/NW/0214 | TAILoR – (Telmisartan and Insulin Resistance in HIV): A Dose-Ranging Phase II Randomised Open-Labelled Trial of Telmisartan as a strategy for the Reduction of Insulin Resistance in HIV-Positive Individuals on Combination Antiretroviral Therapy (cART) | 62 | 31/12/2015 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. Study recruitment window nationally extended from 30/09/2014 to 31/12/2015. |
| C&W14/066 | 14/LO/0803 | Randomized, Placebo-Controlled, Multiple-Dose Study to Evaluate the Pharmacodynamics, Safety and Pharmacokinetics of BMS-955176 (Double-Blinded) and BMS-955176 with Atazanavir +/- Ritonavir (Open-Labelled) in HIV-1 Infected Subjects | 22 | 23/12/2014 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| C&W14/079 | 14/EE/1063 | A Phase III, Open Label, Randomized Study of AZD9291 versus Platinum-Based Doublet Chemotherapy for Patients with Locally Advanced or Metastatic Non-Small Cell Lung Cancer whose Disease has Progressed with Previous Epidermal Growth Factor Receptor Tyrosine Kinase Inhibitor Therapy and whose Tumours harbour a T790M mutation within the Epidermal Growth Factor Receptor Gene | 2 | 31/08/2015 | Open | Not applicable | Trial remains open and therefore is still able to meet recruitment target. |
| HHG09004NI | 09/H1102/54 | An International, Multicentre, Prospective Observational study of the safety of maraviroc used with optimized background therapy in treatment-experienced HIV-1 infected patients | 5 | No date agreed with sponsor | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W10/036 | 10/H0711/34 | A Phase 3, Randomized, Double-Blind Study to Evaluate the Safety and Efficacy of GS 9350-boosted Atazanavir versus Ritonavir-boosted Atazanavir Each Administered with Emtricitabine/Tenofovir Disoproxil Fumarate in HIV 1 Infected, Antiretroviral Treatment-Naive Adults | 11 | 28/02/2013 | Closed - in follow up | Yes | Last patient last visit took place 21/06/2014, with site having met recruitment target. |
| C&W11/044 | 11/LO/0751 | A Phase 3, Open-label Safety study of Cobicistat-containing Highly Active Antiretroviral Regimens in HIV-1 Infected Patients with Mild to Moderate Renal Impairment | 5 | 31/07/2013 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W11/052 | 11/LO/0785 | A Multi-Centre, Randomised, Blinded, Placebo-Controlled Study to Evaluate the Safety of Maraviroc in Combination with Other Antiretroviral Agents in HIV-1 Infected Subjects Co-Infected With Hepatitis C and / or Hepatitis B Virus | 2 | 30/09/2013 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W11/076 | 11/SC/0327 | A Phase 3, Randomised, Double-blind, Placebo-controlled, Parallel-group, Multi-centre Study to Evaluate the Safety and Efficacy of Ustekinumab Induction Therapy in Participants with Moderately to Severely Active Crohn's Disease | 5 | 30/12/2014 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W12/017 | 11/SC/0523 | A Phase 3b Randomized, Open Label Study to Evaluate Switching from Regimens Consisting of a Ritonavirboosted Protease Inhibitor (PI/r) plus Emtricitabine/Tenofovir Fixed-Dose Combination (FTC/TDF) to the Elvitegravir/Cobicistat/Emtricitabine/Tenofovir Disoproxil Fumarate Single-Tablet Regimen (EVG/COBI/FTC/TDF) in Virologically Suppressed, HIV 1 Infected Patients | 8 | 01/03/2013 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W12/018 | 11/SC/0524 | A Phase 3b Randomized, Open-Label Study to Evaluate Switching from Regimens Consisting of a Non-nucleoside Reverse Transcriptase Inhibitor (NNRTI) plus Emtricitabine (FTC) and Tenofovir DF (TDF) to the Elvitegravir/Cobicistat/Emtricitabine/Tenofovir Disoproxil Fumarate Single-Tablet Regimen (EVG/COBI/FTC/TDF) in Virologically-Suppressed, HIV-1 Infected Patients | 8 | 01/03/2013 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W14/063 | 14/LO/0667 | A Phase III Open-Label Clinical Trial to Study the Efficacy and Safety of the Combination Regimen of MK-5172/MK-8742 in Treatment-Naive Subjects with Chronic HCV GT1, GT4, GT5, and GT6 Infection who are Co-Infected with HIV | 9 | 11/05/2015 | Closed - in follow up | No | Trial remained in follow up during this reporting period. 8 patients were recruited, in addition to 5 screen failures. Due to sponsor closing recruitment sooner than anticipated, recruitment of a further 1 patient was not possible. |
| C&W14/062 | 14/SC/0225 | A Phase 3, Randomized, Double-Blind, Switch Study to Evaluate F/TAF in HIV 1 Positive Subjects who are Virologically Suppressed on Regimens containing FTC/TDF (GS-US-311-1089) | 8 | 15/06/2016 | Closed - in follow up | No | Trial remained in follow up during this reporting period. 6 patients were recruited, and due to sponsor closing recruitment sooner than anticipated, recruitment of a further 2 patients was not possible. |
| C&W10/046 | 09/S501/68 | A Randomised Multicenter, Open-Label, Phase 3 Study of Gemcitabine-Cisplatin Chemotherapy Plus IMC-11F8 Versus Gemcitabine-Cisplatin Chemotherapy Alone in the First-Line Treatment of Patients with Squamous Stage IIb or IV Non-Small Cell Lung Cancer (NSCLC) | 4 | 31/01/2013 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W10/103 | 10/H0706/69 | A Phase III, randomised, double blind study of the safety and efficacy of GSK1349572 50 mg once daily to raltegravir 400 mg twice daily both administered with fixed-dose dual nucleoside reverse transcriptase inhibitor therapy over 96 weeks in HIV-1 infected antiretroviral therapy naive adult subjects. | 10 | 31/01/2013 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W12/016 | 11/LO/1974 | A Multicenter, controlled, Open-Label Extension (OLE) Study To Assess the Long-Term Safety and Efficacy of AMG 145 | 2 | 19/03/2013 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W13/016 | 11/LO/1456 | A Long Term Follow-up Registry Study of Subjects Who Did Not Achieve a Sustained Virologic Response in Gilead-Sponsored Trials in Subjects with Chronic Hepatitis C Infection | 5 | 01/10/2016 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W13/039 | 13/LO/0821 | A Phase 3 Open-label Safety Study of Elvitegravir/Cobicistat/Emtricitabine/Tenofovir Alafenamide Single-Tablet Regimen in HIV-1 Positive Patients with Mild to Moderate Renal Impairment (GS-US-292-0112) | 5 | 01/02/2016 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |

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| C&W13/044 | 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 Suppressed, HIV1 Positive Subjects (GS-US-292-0109) | 5 | 22/01/2016 | Closed - in follow up | Yes | Trial remained in follow up during this reporting period. |
| C&W13/050 | 13/LO/0572 | A Phase 3, Randomized, Double-Blind Study to Evaluate the Safety and Efficacy of Elvitegravir/Cobicistat/Emtricitabine/ Tenofovir Alafenamide Versus Elvitegravir/Cobicistat/ Emtricitabine/Tenofovir Disoproxil Fumarate in HIV 1 Positive, Antiretroviral Treatment- Naïve Adults (GS-US-292-0104) | 10 | 22/01/2016 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W13/052 | 13/LO/0574 | A Phase 3, Randomized, Double-Blind Study to Evaluate the Safety and Efficacy of Elvitegravir/Cobicistat/Emtricitabine/ Tenofovir Alafenamide Versus Elvitegravir/Cobicistat/ Emtricitabine/Tenofovir Disoproxil Fumarate in HIV 1 Positive, Antiretroviral Treatment- Naïve Adults (GS-US-292-0111) | 10 | 22/01/2016 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W13/075 | 13/EE/0276 | A Phase 3B Randomized, Open-Label Multi-Center Trial Assessing Sofosbuvir + Ribavirin for 16 or 24 Weeks and Sofosbuvir + Pegylated Interferon + Ribavirin for 12 Weeks in Subjects with Genotype 2 or 3 Chronic HCV Infection (GS-US-334-0153) | 5 | 21/05/2014 | Closed - in follow up | No | Trial remained in follow up during this reporting period. |
| C&W14/087 | 14/LO/1227 | Pharmacokinetics of DOLUTEGRAVIR once daily and ELVITEGRAVIR/COBICISTAT once daily over 10 days following drug intake cessation in healthy volunteers | 16 | 21/01/2015 | Suspended | Not applicable | Trial recruitment has been suspended due to lack of commercial stocks of elvitegravir (to be provided by commercial sponsor). Recruitment will resume once supplies are secure for site. |
| C&W13/026 | 13/LO/0425 | A Randomized, Double-Blind, Controlled Study to Evaluate the Efficacy and Safety of the Combination of ABT-450/Ritonavir/ABT-267 (ABT-450/r/ABT-267) and ABT-333 With and Without Ribavirin (RBV) in Treatment-Naïve Adults with Genotype 1a Chronic Hepatitis C Virus (HCV) Infection (PEARL-IV) | 5 | 01/11/2014 | Closed - follow up complete | No | Last patient last visit took place 29/07/2014, with site not having met recruitment target. |
| C&W10/035 | 10/H0711/33 | A Phase 3, Randomized, Double -Blind Study to Evaluate the Safety and Efficacy of Elvitegravir/Emtricitabine/Tenofovir Disoproxil Fumarate/GS-9350 versus Ritonavir-Boosted Atazanavir Plus Emtricitabine/Tenofovir Disoproxil Fumarate in HIV 1 Infected, Antiretroviral Treatment-Naïve Adults QUAD | 9 | 28/02/2013 | Closed - follow up complete | Yes | Last patient last visit took place 21/06/2014, with site having met recruitment target. |
| C&W12/129 | 12/SC/0540 | A Phase III, Randomised, Partially Double-Blind and Placebo-Controlled Study of BI 207127 in Combination with Faldaprevir and Ribavirin in Treatment-Naïve Patients with Chronic Genotype 1 HCV Infection. | 5 | 01/01/2016 | Closed - follow up complete | Yes | Last patient last visit took place 18/02/2014, with site having met recruitment target. |
| C&W13/013 | 13/LO/0006 | A Phase 3, Open-label Study to Investigate the Efficacy and Safety of Sofosbuvir plus Ribavirin in Chronic Genotype 1, 2, 3 and 4 Hepatitis C Virus (HCV) and Human Immunodeficiency Virus (HIV) Co-infected Subjects | 5 | 15/07/2014 | Closed - follow up complete | Yes | Last patient last visit took place 01/07/2014, with site having met recruitment target. |
| C&W12/075 | 12/NE/0266 | A Phase 3, Multicenter, Randomized, Double-Blind, Placebo-Controlled Study to Investigate the Efficacy and Safety of GS-7977 + Ribavirin for 12 Weeks in Treatment Naïve and Treatment Experienced Subjects with Chronic Genotype 2 or 3 HCV Infection | 5 | 01/10/2014 | Closed - follow up complete | Yes | Last patient last visit took place 08/01/2014, with site having met recruitment target. |
| C&W13/010 | 12/EE/0400 | An Open-Label Study of GS-7977+ Ribavirin for 12 Weeks in Subjects with Chronic HCV Infection who Participated in Prior Studies Evaluating GS-7977 | 1 | 04/09/2014 | Closed - follow up complete | Yes | Last patient last visit took place 11/02/2014, with site having met recruitment target. |
| C&W12/047 | 12/LO/0497 | Multicenter, Open-Label Study of Telaprevir in Combination With Peginterferon Alfa and Ribavirin in Human Immunodeficiency Virus/Genotype 1 Chronic Hepatitis C Coinfected Subjects With Severe Fibrosis or Compensated Cirrhosis | 3 | 01/02/2014 | Closed - follow up complete | No | Last patient last visit took place 22/04/2014, with site not having met recruitment target. |
| C&W11/018 | 11/SC/0007 | A phase III, randomised, double-blind and placebo-controlled study of once daily BI 201335 120mg for 12 or 24 weeks or BI 201335 240mg for 12 weeks in combination with pegylated interferon-a and ribavirin in treatment-naïve patients with genotype 1 chronic hepatitis C infection | 8 | 28/02/2014 | Closed - follow up complete | No | Last patient last visit took place 21/11/2013, with site having not met recruitment target. |
| C&W13/022 | 13/LO/0129 | A Phase 4 Cross-Sectional Study of Bone Mineral Density in HIV-1 Infected Subjects (GS-US-104-0423) | 6 | 12/07/2014 | Closed - follow up complete | No | Last patient last visit took place 10/03/2014, with site having not met recruitment target. This was a roll over study, to which 4 patients were recruited. Due to eligibility being based upon participation in the original study, recruitment of a further 2 patients was not possible. |
| C&W13/057 | 13/LO/0830 | Phase 3 open label study evaluating the efficacy and safety of pegylated interferon lambda-1a, in combination with ribavirin and daclatasvir, for treatment of chronic HCV infection with treatment naïve genotypes 1, 2, 3 or 4 in subjects co-infected with HIV | 5 | 15/02/2015 | Closed - follow up complete | No | Last patient last visit took place 29/09/2014, with site having not met recruitment target. 4 patients were recruited, and due to sponsor closing recruitment sooner than anticipated, recruitment of a further 1 patient was not possible. |
| C&W13/073 | 13/LO/1290 | A Follow-up Study to Assess Resistance and Durability of Response to AbbVie Direct-Acting Antiviral Agent (DAA) Therapy in Subjects Who Participated in Phase 2 or 3 Clinical Studies for the Treatment of Chronic Hepatitis C Virus (HCV) Infection | 3 | 08/11/2016 | Withdrawn | No | Trial closed by sponsor due to a change in development pipeline within sponsor company. Trial is a roll over trial, and sponsor decided to close the trial on 02/04/2014, prior to any patients rolling over on to the trial at site. As such, sponsor did not expect any recruitment at site. |