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Project Name: GS-US-248-0132 Form: 12-Lead ECG [ECG12]

Was an ECG performed?	Yes No
If No, please comment why ECG was not done:	
Date of ECG:	/ /
Time of ECG:	:
HR Rate (bpm):	
PR Interval (msec):	
QRS Interval (msec):	
QT Interval (msec):	
QTcF Interval (msec) (Calculated)	
Overall Assessment of ECG:	Normal Abnormal
If Abnormal, please describe the abnormality:	
Significance of Abnormality:	Not Clinically Significant  Clinically Significant

Project Name: GS-US-248-0132	
Form: Adverse Event Summary [AES]	
Generated On: Aug-01-2013 06:49:06	
Did the subject experience any Adverse Events during the course of the study?	Yes
	No

Form: Adverse Event [AE] Generated On: Aug-01-2013 06:49:06 Adverse Event: (List diagnosis or each symptom separately) AE serious: Yes (If the Adverse Event is Serious, please click the [?] help text to open the SAE Report Form in another window. Print and fill in all the information for the first three pages on the SAE form and fax to the appropriate CRO contact) Start Date: (DD-MMM-YYYY) End Date: (DD-MMM-YYYY) Please check if Ongoing: Related to any of the Study Drugs: Yes (All study drugs are to be considered in assessment of relationship to study drugs: No GS-5885, GS-9451, Tegobuvir/Placebo, RBV/Placebo) Related to Study Procedures: Yes No Study Drug Action Taken - GS-5885: No change Interrupted Discontinued Dose Reduced Study Drug Action Taken - GS-9451: No change Interrupted Discontinued Dose Reduced Study Drug Action Taken - Tegobuvir/Placebo: No change Interrupted

Project Name: GS-US-248-0132

Discontinued

Dose Reduced

Generated On: Aug-01-2013 06:49:06 Study Drug Action Taken - RBV/Placebo: No change Interrupted Discontinued Dose Reduced Severity: Mild Moderate Severe Life-Threatening Other Action Taken: (Check all that apply) None: Medication Required: Other Treatment Required: Hospitalized/Prolonged Hospitalization: Hidden field for AE\_AESER\_ALERT Note: To add additional adverse events, please complete an additional form, by clicking on 'Add a new log line'. If Adverse Event is Serious, please refer to the regulatory binder for an SAE Report form. Complete an SAE Report and E-mail or fax the SAE Report within 24 hours of the Reporter's knowledge of the event. Contact information for faxing and e-mailing this form is found in the study protocol in section 8.5.1. A Serious Adverse Event is any adverse experience that results in any of the following outcomes: (1) Death, (2) Is life-threatening (at immediate risk of death at time of the event), (3) Requires subject hospitalization or prolonged hospitalization, (4) Persistent or significant disability/incapacity, (5) Congenital abnormality/birth defect or (6) Any other important Medical Event based on clinician's judgment or may require medical or surgical intervention to prevent one of the other serious criteria.

Refer to protocol for protocol-specific SAE definitions.

Project Name: GS-US-248-0132

Form: Adverse Event [AE]

Form: Complete Physical Examination [PE] Generated On: Aug-01-2013 06:49:06

Select a response for each body system. If a body system is not examined, select "Not Done."

Body System	Head, Neck & Thyroid	
	Eyes, Ears, Nose, Throat, Mouth & Tongue	
	Chest (Excluding breasts)	
	Respiratory	
	Cardiovascular	
	Lymph Nodes	
	Abdomen	
	Skin, Nails & Hair	
	Musculoskeletal	
	Neurological	
	Retinal Exam Results	
	Other	
If Body System is Other, please specify:		
Result:	Abnormal	
	Normal	
	Not Done	
If "Abnormal", please describe abnormal physical findings:		
If additional Body Systems are needed then click on 'Add New Log Line' at the bottom of the form.		

Project Name: GS-US-248-0132		
Form: Concomitant Medication Summary [CMS]		
Generated On: Aug-01-2013 06:49:06		
Did the subject take any Medications from time of informed consent signing through study completion?	Yes No	

Project Name: GS-US-248-0132 Form: Concomitant Medication [CM] Generated On: Aug-01-2013 06:49:06

Drug Name:		
Indication: (Use Medical Terminology)		
Route:	Inhalation  Intradermal  Intralesional  Intramuscular  Intranasal  Intravenous  Intraocular  Intraperitoneal  Oral  Rectal  Subcutaneous  Sublingual  Topical  Vaginal	
If "Other", places specify:	Other	
If "Other", please specify:		
Start Date: (DD-MMM-YYYY)	1 1	
Stop Date: (DD-MMM-YYYY)	/ /	
Please check if Ongoing:		
Note: To add additional concomitant medications, please complete an additional form, by clicking on 'Add a new log line'.		

Project Name: GS-US-248-0132 Form: Death Report [DEATH]

Immediate Cause of Death:

Generated On: Aug-01-2013 06:49:06

In addition to this eCRF, please refer to the regulatory binder for an SAE Report form. Complete an SAE Report and E-mail or fax the SAE Report within 24 hours of the Reporter's knowledge of the event. Please also send a copy of the discharge/death summary and autopsy report, if available.

Contact information for faxing and e-mailing these forms is found in the study protocol in section 8.5.1(Click the [?] help text to open the SAE Report Form in another window. Please fill in all appropriate information on the first three pages before faxing the forms to the appropriate CRO contact.)

Please update the Adverse Event eCRF, as needed.

Date of Death: (DD-MMM-YYYY)

Form: Demographics [DM] Generated On: Aug-01-2013 06:49:06 Date of Birth: (DD-MMM-YYYY) Age: Sex: Male Female

Ethnicity: Hispanic or Latino Not Hispanic or Latino Not Permitted Race: American Indian or Alaska Native Asian Black or African Heritage Native Hawaiian or Pacific Islander White Not Permitted Other If "Other", please specify: Year of Birth (yyyy) / /

Project Name: GS-US-248-0132

Form: Dose Administration (Day 1) [PKEX] Generated On: Aug-01-2013 06:49:06

Name of Actual Treatment:	GS-5885 GS-9451 Tegobuvir/Placebo RBV/Placebo
Date of First Dose: (DD-MMM-YYYY)	/ /
Time of First Dose: (00:00-23:59)	
Dose:	
Dose Units:	mg ug tablets capsules
Dose Taken With Food	Yes No

Form: Dose Administration (Serial PK) [PKEX2]

Dose Timepoint:	Pre-PK Day Dose
	$\equiv$
	PK Day AM Dose
	PK Day PM Dose
	Post-PK Day AM Dose
Name of Actual Treatment:	GS-5885
	GS-9451
	Tegobuvir/Placebo
	RBV/Placebo
Date of Dose: (DD-MMM-YYYY)	/ /
Time of Dose: (00:00-23:59)	:
Dose:	
Dose Units:	mg
	ug
	tablets
	capsules
Dose Taken With Food:	Yes
	No

Form: Dose Administration (Single PK) [PKEX1]

Check 'Not Applicable' box if Subject is participating in Serial PK Substudy at this visit:	
Name of Actual Treatment:	GS-5885 GS-9451 Tegobuvir/Placebo RBV/Placebo
Date of Dose prior to Sample Draw: (DD-MMM-YYYY)	1 1
Time of Dose prior to Sample Draw: (00:00-23:59)	
Dose:	
Dose Units:	mg ug tablets capsules
Dose Taken With Food	Yes No

Form: Dose Administration (Viral Dynamic Substudy) [PKEX3]

Name of Actual Treatment:	GS-5885 GS-9451 Tegobuvir/Placebo RBV/Placebo
Date of Dose prior to Sample Draw: (DD-MMM-YYYY)	/ /
Time of Dose prior to Sample Draw: (00:00-23:59)	
Dose:	
Dose Units:	mg ug tablets capsules
Dose Taken With Food:	Yes No

Project Name: GS-US-248-0132 Form: General Comments [CO]

Visit:	Screening
	Initial Treatment Visits
	Baseline/Day 1
	Week 1
	Week 2
	Week 4
	Week 6
	Week 8
	Week 10
	Week 12
	Week 16
	Week 20
	Week 24
	Early Termination (Initial Treatment)
	Study Drug Completion (Initial Treatment)
	PK and/or Viral Dynamic Substudies
	Day 1
	Day 2
	Day 3
	Day 4
	Day 5
	Day 10
	PK Week 2
	Post-Treatment Follow-Up Visits
	Post-Treatment Follow-Up Week
	Post-Treatment Follow-Up Week 8
	Post-Treatment Follow-Up Week
	Post-Treatment Follow-Up Week
	Post-Treatment Follow-Up Week 20

Project Name: GS-US-248-0132	
Form: General Comments [CO]	
Generated On: Aug-01-2013 06:49:06	
	Post-Treatment Follow-Up Week
	Post-Treatment Follow-Up Month 6
	Adverse Events
	Concomitant Medication
	Study Drug Administration
	Study Drug Accountability
	Pregnancy Report
	Death
	General Comments
	Unscheduled
	Study Completion
	Missed Dose Overdose Log

Project Name: GS-US-248-0132 Form: General Comments [CO]

Form:	Adverse Event
	Adverse Event Summary
	Complete Physical Examination
	Concomitant Medication
	Concomitant Medication Summary
	Death Report
	Demographics
	Dose Administration (Day 1)
	Dose Administration (Single PK )
	Dose Administration (Serial PK)
	Dose Administration (Viral Dynamic Substudy)
	General Comments
	Inclusion/Exclusion Criteria
	Investigator's Signature
	Liver Fibrosis Assessment
	Medical History
	Pregnancy Report
	Randomization
	Single PK Plasma
	Serial PK Plasma
	PK Plasma (Viral Dynamic Substudy Day 1)
	Stop Treatment
	Study Completion
	Study Drug Accountability - GS- 5885
	Study Drug Accountability - GS- 9451
	Study Drug Accountability - Tegobuvir//Placebo
	Study Drug Accountability - RBV
	Study Drug Administration - Initial Treatment
	Study Drug Completion (Initial Treatment)

Generated On: Aug-01-2013 06:49:06 Subject Follow-Up Status Urine Pregnancy Test Urine Pregnancy Test (Baseline) Visit Date 12-Lead ECG Missed Dose and Overdose Log - Initial Treatment Prior HCV Treatment History Interferon Classification Interferon Ineligibility Reasons Interferon Intolerant Reasons Vital Signs Vital Signs (Screening) Vital Signs with Weight General Comments: General Comments (Continued) Note: To add additional comments, please complete an additional form, by clicking on 'Add a new log line'.

Project Name: GS-US-248-0132 Form: General Comments [CO]

Form: Inclusion/Exclusion Criteria [IE]
Generated On: Aug-01-2013 06:49:06

Date informed Consent Signed: (DD-MMM-YYYY)

Under which protocol version was the subject enrolled?

Original
Amendment 1

Was subject a Screen Failure?

Pes
No

No

If "No", Please complete the following (Please tick all Inclusion criteria not met or Exclusion criteria met):

Project Name: GS-US-248-0132

Project Name: GS-US-248-0132
Form: Inclusion/Exclusion Criteria [IE]
Generated On: Aug-01-2013 06:49:06
Inclusion/Exclusion criteria:

Inclusion/Exclusion criteria:		Inclusion 1
		Inclusion 2
		Inclusion 3
		Inclusion 4
		Inclusion 5
(		Inclusion 6
		Inclusion 7
		Inclusion 8
	$\supseteq$	Inclusion 9
	$\exists$	Inclusion 10
	ᆜ	Inclusion 11
	ᆜ	Inclusion 12
	ᆜ	Inclusion 13
	ᆜ	Inclusion 14
ļ	ᆜ	Inclusion 15
	ᆜ	Inclusion 16
	ᆜ	Inclusion 17
ļ	ᆜ	Exclusion 1
l	닉	Exclusion 2
	닉	Exclusion 3
	닉	Exclusion 4
	닉	Exclusion 5
l	닉	Exclusion 6
	닉	Exclusion 7
l de la companya de	닉	Exclusion 8
	닉	Exclusion 9
	닉	Exclusion 10
	닉	Exclusion 11
		Exclusion 12

Form: Inclusion/Exclusion Criteria [IE] Generated On: Aug-01-2013 06:49:06 Exclusion 13 Exclusion 14 Exclusion 15 Exclusion 16 Exclusion 17 Exclusion 18 Exclusion 19 Exclusion 20 Exclusion 21 Exclusion 22 Exclusion 23 Exclusion 24 Exclusion 25 Exclusion 26 Exclusion 27 Exclusion 28 Exclusion 29 Inclusion criteria not met/Exclusion criteria met? If subject that did not meet inclusion/exclusion criteria was enrolled, please provide an explanation. For subjects who are screen failures but meet eligibility criteria - ONLY: Adverse Event Provide the most significant reason why the subject was not randomized in the IWRS. Investigator Decision Subject Withdrew Consent Lost to Follow-Up Outside of Visit Window Study Enrollment Closed Other

Project Name: GS-US-248-0132

Project Name: GS-US-248-0132	
Form: Inclusion/Exclusion Criteria [IE]	
Generated On: Aug-01-2013 06:49:06	
If "Other", specify:	

Project Name: GS-US-248-0132	
Form: Interferon Classification [INTC]	
Generated On: Aug-01-2013 06:49:07	
Please indicate if subject is interferon ineligible or interferon intolerant	Interferon ineligible
	Interferon intolerant

Form: Interferon Ineligibility Reasons [INTIR]

Generated On: Aug-01-2013 06:49:07

Please choose the primary Interferon Ineligibility Reason as per Inclusion Criteria #6.

If additional secondary reasons are present, please click on the 'Add a new log line' link at the bottom of the table below.		
Primary/Secondary Reasons	Primary Reason	
	Secondary Reason(s)	
Interferon Ineligibility Reasons as per Inclusion Criteria #6	Autoimmune Disorder	
	Significant Psychiatric Disease	
	Seizure Disorder	
	Thyroid Dysfunction	
	Retinal Disease	
	Poorly Controlled Diabetes	
	Other (as approved by the Medical Monitor)	
Interferon Ineligibility Reasons Specify		
Date of Diagnosis:(DD-MMM-YYYY)	/ /	

Form: Interferon Intolerant Reasons [INTIAE]

Generated On: Aug-01-2013 06:49:07

Please choose the primary Interferon Intolerant Reason as per Inclusion Criteria #6.

If additional secondary reasons are present, please click on the 'Add a new log line' link at the bottom of the table below.	
Primary/Secondary Reasons	Primary Reason
	Secondary Reason(s)
Interferon Intolerant Reasons as per Inclusion Criteria #6	Significant Local or Systemic Adverse Reaction
	Psychiatric Disease
	Significant Cognitive Impairment
	Neuropathy
	Disabling Flu-Like Symptoms
	Gastrointestinal Toxicity
	Thrombocytopenia
	Neutropenia
	Retinal Disease
	Autoimmune Disorder
	Other (as approved by the Medical Monitor)
Interferon Intolerant Reason Specify	
Date of Onset	/ /

Project Name: GS-US-248-0132	
Form: Investigator's Signature [INVSIG]	
Generated On: Aug-01-2013 06:49:07	
By entering my Medidata password, I affirm that I have reviewed and evaluated the case reportforms and verify that they accurately reflect the information in the source documents for thissubject. I understand source documentation can include (but is not limited to) medical records, laboratory results, x-rays, electronic communications, etc.	

Form: Liver Fibrosis Assessment [BIOPSY] Generated On: Aug-01-2013 06:49:07

LIVER BIOPSY	
Check if Biopsy not done:	
Date of Liver Biopsy: (DD-MMM-YYYY)	1 1
Fibrosis Staging Method:	Metavir
	Ishak
	Knodell
	Batts-Ludwig
	Scheuer
Fibrosis Staging Result:	F0
	F0-F1
	F1
	F1-F2
	F2
	F2-F3
	F3
	F3-F4
	F4
	F5
	F6
Cirrhosis:	Yes
	No
NON-INVASIVE ALTERNATIVE TO LIVER BIOPSY	
Check if not done:	
Date of Procedure: (DD-MMM-YYYY)	/ /

Generated On: Aug-01-2013 06:49:07

Method:

FibroTest
FibroScan
Acoustic Radiation Force Impulse Imaging
Other

If Method is Other, please specify:

Result:

Cirrhosis

Yes

No

Project Name: GS-US-248-0132

Form: Liver Fibrosis Assessment [BIOPSY]

Project Name: GS-US-248-0132 Form: Medical History [MH]

Generated On: Aug-01-2013 06:49:07

Please record date of diagnosis for Hepatitis C on row 1 and indicate if ongoing by checking the 'Ongoing' box.

In addition, click on the 'Add a new Log line' link below	and add any relevant medical history.
Condition:	
Start Date: (DD-MMM-YYYY)	/ /
Stop Date: (DD-MMM-YYYY)	/ /
Check if Condition is Ongoing:	
Note: To add additional Medical History entries, please of line'.	complete an additional form, by clicking on 'Add a new log

Form: Missed Dose and Overdose Log [EXMSOV]

Generated On: Aug-01-2013 06:49:07

Only enter a record for days on which the dairy indicates a missed dose or overdose of a study drug. Enter all dosing information for that day.

If a subject mistook an evening dose for the QD drugs GS-9451 or GS-5885 and this was indicated in the dairy, please enter them in the last two columns. If an evening dose of GS-9451 or GS-5885 were not taken, please leave the column blank.

Date (dd-mmm-yyyy)	/ /
Morning Dosing:	
Please record number of tablets taken: Ribavirin	
Morning Dosing:	
Please record number of capsules taken: <b>Tegobuvir</b>	
Morning Dosing:	
Please record number of tablets taken: GS-9451	
Morning Dosing:	
Please record number of tablets taken: GS-5885	
Evening Dosing:	
Please record number of tablets taken: <b>Ribavirin</b>	
Evening Dosing:	
Please record number of capsules taken: <b>Tegobuvir</b>	
Evening Dosing:	
Please record number of tablets taken: GS-9451	
Evening Dosing:	
Please record number of tablets taken: GS-5885	
If and overdose of study drug has been reported, please click on the [?] help text to open the Oversdose Form in	
another window. Fax or email the completed form to the appropriate CRO contact.	

Form: PK Plasma (Viral Dynamic Substudy Day 1) [PKS1]
Generated On: Aug-01-2013 06:49:07

Will a serial PK profile be collected for this subject?

Yes

No

IF YES, COMPLETE THE REMAINDER OF THIS FORM

Date of PK Collection: (DD-MMM-YYYY)

Dose Time Point:

Collection Time 4.0 Hours Post-dose
Collection Time 6.0 Hours Post-dose
Collection Time 8.0 Hours Post-dose
Time of PK Collection: (00:00-23:59)

Time of PK Collection: (00:00-23:59)

Please specify why this timepoint was not collected:

Project Name: GS-US-248-0132
Form: Pregnancy Report [PREGREP]
Generated On: Aug-01-2013 06:49:07

f a pregnancy occurs, please complete as mu

If a pregnancy occurs, please complete as much information as possible on this form.

In addition to this eCRF, refer to the regulatory binder and complete the 'Pregnancy Report' and email or fax the report within 24 hours of the Reporter's knowledge of the event. When the outcome of the pregnancy is known, complete the 'Pregnancy Outcome Report'.(Click the [?] help text to open the Pregnancy Report Form in another window. Please fill in all information before faxing the forms to the appropriate CRO contact.)

Contact information for faxing and e-mailing these forms to the CRO during the conduct of the study is found in the study protocol in section 8.5.1. Additional reporting details may be found in protocol section 8.7.

Please also report the pregnancy to the Ribavirin Pregnancy Registry at ribavirinpregnancyregistry.com

Last Menstrual Period: (DD-MMM-YYYY)

// /

Pregnancy Confirmed: (DD-MMM-YYYY)

(Click the [?] help text to open the Pregnancy Outcome Report Form in another window. Please fill in all information on the first 3 pages before faxing the forms to the appropriate CRO contact.)

Form: Prior HCV Treatment History [PRHCV2]

HCV Treatment:  HCV treatment Specify	Interferon-alfa Pegylated Interferon-alfa Other Interferon (specify) Ribavirin Other HCV treatment (specify)
Starting Dose:	
Dose Unit:	Ug Ug/kg Mg Other
Dose Unit Specify	
Dose Unit Specify  Frequency	Once a Week Three Times a Day Once a Day Twice a Week Other
	Three Times a Day Once a Day Twice a Week
Frequency	Three Times a Day Once a Day Twice a Week
Frequency Frequecy Specify	Three Times a Day Once a Day Twice a Week Other

Project Name: GS-US-248-0132
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Form: Prior HCV Treatment History [PRHCV2]

Generated On: Aug-01-2013 06:49:07

Stop	Year	(yyyy)

/ /

Project Name: GS-US-248-0132 Form: Randomization [RAND]

Note: All baseline tests and procedures, including ECGs, must be completed prior to randomization and dosing/dispensing of GS-5885, GS-9451, Tegobuvir/Placebo or RBV/Placebo.		
Was the Subject Randomized?	Yes No	
4-Digit Subject Number as assigned by IWR system		
Will the subject participate in the PK Substudy?	Yes No	
Date of PK Substudy Consent (DD-MMM-YYYY)	/ /	
Will the subject participate in the Viral Dynamic Substudy?	Yes No	
Date of Viral Dynamic Substudy Consent: (DD-MMM-YYYY)	/ /	
Will the subject participate in Pharmacogenomic Testing?	Yes No	
Date of Pharmacogenomic Testing Consent: (DD-MMM-YYYY)	/ /	
RANDNUM+Z_SUBID		

Form: Registry Substudy [REGISTRY] Generated On: Aug-01-2013 06:49:07 Which Registry study is the subject eligible to enroll in? None SVR Registry (248-0122) Sequence Registry (248-0123) NA, Site Not Participating Will the subject enroll in the registry study? Yes No If No, state the primary reason why the subject will not enroll? Subject is not interested in participating Subject will receive other HCV treatment Lost to follow-up

Project Name: GS-US-248-0132

Other, specify

Other

Project Name: GS-US-248-0132 Form: Serial PK Plasma [PKS]

Will a serial PK profile be collected for this subject?	Yes No
IF YES, COMPLETE THE REMAINDER OF THIS FORM	
Date of PK Collection: (DD-MMM-YYYY)	/ /
PK Time Point:	Collection Time Pre-dose  Collection Time 1.0 Hour Post-dose  Collection Time 2.0 Hours Post-dose  Collection Time 3.0 Hours Post-dose  Collection Time 4.0 Hours Post-dose  Collection Time 6.0 Hours Post-dose  Collection Time 8.0 Hours Post-dose  Collection Time 10.0 Hours Post-dose  Collection Time 12.0 Hours Post-dose  Collection Time 12.0 Hours Post-dose  Collection Time 24.0 Post-dose
Time of PK Collection: (00:00-23:59)	:
Not Collected	
Please specify why this timepoint was not collected:	
Collection Time 24.0 Hours Post-Dose: Not Collected	
Collection Time 24.0 Hours Post-Dose: Date of PK Collection: (DD-MMM-YYYY)	/ /
Collection Time 24.0 Hours Post-Dose: Time of PK Collection: (00:00-23:59)	
Collection Time 24.0 Hours Post-Dose: Please specify why this timepoint was not collected:	

Form: Single PK Plasma [PK]
Generated On: Aug-01-2013 06:49:07

Was Single PK Plasma Sample collected?

Pes
No

Date of PK Collection: (DD-MMM-YYYY)

Time of PK Collection: (00:00-23:59)

Generated On: Aug-01-2013 06:49:07

Did the subject complete the protocol-planned duration of the study?

Yes

No

No

Adverse Event

Death

Pregnancy

Efficacy Failure

Protocol Violation

Subject Withdrew Consent

Lost to Follow-Up

Investigator Decision

Study Discontinued by Sponsor

Project Name: GS-US-248-0132

Form: Study Completion [STUDCOMP]

Form: Study Drug Accountability - GS-5885 [DA1]

Visit Dispensed:	Baseline
	Week 4
	Week 8
	Week 12
	Week 16
	Week 20
	Unscheduled
Drug Name:	GS-5885
	GS-9451
	RBV/Placebo
	Tegobuvir/Placebo
	GS-5885 Rescue Therapy
	GS-9451 Rescue Therapy
	RBV Rescue Therapy
	Tegobuvir/Placebo Rescue Therapy
	PEG Rescue Therapy
Date Dispensed:	/ /
5-Digit Bottle Number:	
Number of Tablets Dispensed:	
Date Returned:	/ /
Number of Tablets Returned:	
Study Drug Not Returned:	

Form: Study Drug Accountability - GS-9451 [DA2]

Visit Dispensed:	Baseline
	Week 4
	Week 8
	Week 12
	Week 16
	Week 20
	Unscheduled
Drug Name:	GS-5885
	GS-9451
	RBV/Placebo
	Tegobuvir/Placebo
	GS-5885 Rescue Therapy
	GS-9451 Rescue Therapy
	RBV Rescue Therapy
	Tegobuvir/Placebo Rescue Therapy
	PEG Rescue Therapy
Date Dispensed:	/ /
5-Digit Bottle Number:	
Number of Tablets Dispensed:	
Date Returned:	/ /
Number of Tablets Returned:	
Study Drug Not Returned:	

Form: Study Drug Accountability - RBV/Placebo [DA4]

Visit Dispensed:	Baseline
	Week 4
	Week 8
	Week 12
	Week 16
	Week 20
	Unscheduled
Drug Name:	GS-5885
	GS-9451
	RBV/Placebo
	Tegobuvir/Placebo
	GS-5885 Rescue Therapy
	GS-9451 Rescue Therapy
	RBV Rescue Therapy
	Tegobuvir/Placebo Rescue Therapy
	PEG Rescue Therapy
Date Dispensed:	/ /
5-Digit Bottle Number:	
Number of Tablets Dispensed:	
Date Returned:	/ /
Number of Tablets Returned:	
Study Drug Not Returned:	

Form: Study Drug Accountability - Tegobuvir/Placebo [DA3]

Visit Dispensed:	Baseline
	Week 4
	Week 8
	Week 12
	Week 16
	Week 20
	Unscheduled
Drug Name:	GS-5885
	GS-9451
	RBV/Placebo
	Tegobuvir/Placebo
	GS-5885 Rescue Therapy
	GS-9451 Rescue Therapy
	RBV Rescue Therapy
	Tegobuvir/Placebo Rescue Therapy
	PEG Rescue Therapy
Date Dispensed:	/ /
5-Digit Bottle Number:	
Number of Capsules Dispensed:	
Date Returned:	/ /
Number of Capsules Returned:	
Study Drug Not Returned:	

Project Name: GS-US-248-0132
Form: Study Drug Administration (

Form: Study Drug Administration [EX] Generated On: Aug-01-2013 06:49:07

Drug Name:	GS-5885
	GS-9451
	Tegobuvir/Placebo
	RBV/Placebo
Dose:	
Dose Units / Frequency:	capsules/day
	tablets/day
	Other
If "Other", please specify:	
Start Date: (DD-MMM-YYYY)	/ /
Stop Date: (DD-MMM-YYYY)	/ /
Ongoing:	
Check box if study drug was permanently discontinued:	

Generated On: Aug-01-2013 06:49:07

Did subject complete study drug treatment through Week 24?

Yes

No

No

If "No", please specify reason for study drug discontinuation:

Adverse Event

Death

Pregnancy

Efficacy Failure

Protocol Violation

Subject Withdrew Consent

Lost to Follow-Up

Investigator Decision

Randomized but Subject Never
Dosed with Study Drug

Study Discontinued by Sponsor

Project Name: GS-US-248-0132

Form: Study Drug Completion [SDRGCOMP]

Project Name: GS-US-248-0132	
Form: Subject Follow-Up Status [DS]	
Generated On: Aug-01-2013 06:49:07	
Provide the subject status:	
If subject has discontinued, please complete the Study Completion form. For female subjects of childbearing potential, please also continue to complete the follow-up Urine Pregnancy Tests until Month 6	

Form: Subject Number [SUBID] Generated On: Aug-01-2013 06:49:07	
Screening Number (XXX):	
Subject Initials (XXX):	
SCRNID + SUBJINIT	

Project Name: GS-US-248-0132	
Form: Urine Pregnancy Test (Baseline) [PREGTEST1]	
Generated On: Aug-01-2013 06:49:07	
Is female subject of childbearing potential?	Yes
	No
Date Test Performed: (DD-MMM-YYYY)	/ /
Test Result:	Negative
	Positive
	Not Done
Note: A positive urine pregnancy test must be immediately confirmed with a serum pregnancy test, and the subject	
must not be enrolled.	

Project Name: GS-US-248-0132	
Form: Urine Pregnancy Test [PREGTEST]	
Generated On: Aug-01-2013 06:49:07	
Is female subject of childbearing potential?	Yes
	No
Date Test Performed: (DD-MMM-YYYY)	/ /
Test Result:	Negative
	Positive
	Not Done
Note: A positive urine pregnancy test must be immediately confirmed with a serum pregnancy test	

Form: Visit Date [VISDT] Generated On: Aug-01-2013 06:49:07 Was Post-Treatment Follow-Up Week 4 visit performed? Yes No Visit Date: (DD-MMM-YYYY) Was visit performed on the same day as Week 1 Yes No Reason for Unscheduled Visit: (Check all that apply) Evaluation of AE and/or Concomitant Medications: Vital Signs: Physical Exam: ECG: PK: Lab Evaluations: Study Drug Adjustment or Dispensation: Other: Other, Specify:

Project Name: GS-US-248-0132 Form: Vital Signs (Screening) [VS1] Generated On: Aug-01-2013 06:49:08

Weight:	kg/ lb
Height:	cm/ in
Blood Pressure Systolic: (mmHg)	
Blood Pressure Diastolic: (mmHg)	
Pulse (beats/minute):	
Respiration: (breaths/minute)	
Temperature:	Celsius/ Fahrenheit

Project Name: GS-US-248-0132 Form: Vital Signs with Weight [VS2] Generated On: Aug-01-2013 06:49:08

Weight:	kg/ lb
Blood Pressure Systolic: (mmHg)	
Blood Pressure Diastolic: (mmHg)	
Pulse (beats/minute):	
Respiration: (breaths/minute)	
Temperature:	Celsius/ Fahrenheit

Form: Vital Signs [VS] Generated On: Aug-01-2013 06:49:08	
Blood Pressure Systolic: (mmHg)	
Blood Pressure Diastolic: (mmHg)	
Pulse (beats/minute):	
Respiration: (breaths/minute)	
Temperature:	Celsius/ Fahrenheit