

Screening

Form		Keys	Description
SE	Screening Evaluation	ID, RSC	Initial screening form including demographics, baseline physical exam, baseline medications, symptoms, serologies, and labs
SC	Screening Criteria	ID, RSC	Checklist containing all inclusion/exclusion criteria to determine study eligibility.
PE	Pathology Evaluation	ID	Pathology data as completed by central pathologist. Section II is all initial scoring. Section III is additional scoring appended to the database in December 2004.
RA	Risk Assessment	ID	Obtains information on known risk factors associated with Hepatitis C, PI determination of infection source, and date of infection.

Clinic Assessments

Form		Keys	Description
		,	Patient evaluation during the treatment period
			including physical exam, patient reported symptoms,
TE	Treatment Evaluation	ID, TMPT	concomitant medications, depression, and most recent dose of study medications
	Trodition Evaluation	10, 11111	Patient evaluation during the follow-up period
			including physical exam, patient reported symptoms,
			concomitant medications, depression, and most
FE	Follow-up Evaluation	ID, TMPT	recent dose of study medication (only provided at follow-up week 4 visit).
	r onotr up Evaluation	15, 111111	Patient evaluation if patient was seen at anytime
			during the study period outside of protocol timepoints
UE	Unscheduled Visit	ID DOEDATE	including physical exam, patient reported symptoms, concomitant medications, and depression.
UE	Offscrieduled visit	ID, DOEDATE	Checklist of specified educational topics to be
			covered by the coordinator with the patient at the
			baseline visit. Captures date and time of first doses
MA	MATI -HepC	ID	of peg-interferon and ribavirin and prescribed dose of study medications.
IVIA	WATT-Hepc	טון	Checklist of specified educational topics to be
			covered by the coordinator with the patient at
MB	Brief MATI	ID, TMPT	specified timepoints during the treatment period.
LE	Laboratory Evaluation	ID, TMPT, DSMDATE	Captures all laboratory tests completed throughout the study.
	Laboratory Lvaldation	ID, TIVIT I, DOIVIDATE	tilo stady.



Patient Questionnaires

Form		Keys	Description
SQ	Screening Questionnaire	ID	Baseline patient information including demographics, education and work status, alcohol, and smoking history.
QL	Quality of Life	ID, TMPT	Questionnaire based on the SF-36 to measure the patient's quality of life through questions about the patient's physical activities and emotional health.
SS	Social Support and Self-efficacy	ID, TMPT	Designed to measure the amount of social support the patient receives and the patient's level of confidence in performing certain activities.
CD	CES-D	ID, TMPT	CES-D questionnaire which measures the level of depression at specified timepoints during the study.
SA	Symptom Assessment	ID, TMPT	Captures specific symptoms on a continuous VAS line (10 cm). Measurement is the distance, in centimeters, from the left axis to the point where the line was marked by the participant.
XM	Sexual Function: Male	ID, TMPT	Measures sexual functioning in males at specified timepoints throughout the study.
XF	Sexual Function: Female	ID, TMPT	Measures sexual functioning in females at specified timepoints throughout the study.
AD	Adherence Questionnaire	ID, TMPT	Questionnaire designed to monitor patient's adherence to study medications during the treatment period, e.g. study medication taking behavior, missed doses, and reasons for missed doses.
MM	MEMS Questionnaire	ID, TMPT	Questionnaire designed to monitor the patient's adherence to the study medication schedule through use of the MEMS cap.

Other Study Forms

Form		Keys	Description
TW	Trackcap Worksheet	ID	Records all refill and non-event openings for distributed MEMS caps for each patient.
AE	Adverse Events	ID, PNUM, LNUM	Obtains information regarding adverse events that occurred during the study period.
DC	Dose Change Form	ID, PNUM, LNUM	Tracks physician prescribed changes to study medication dosages throughout the treatment period.
DF	Discontinuation Form	ID, DDCDATE	Documents patient discontinuation from treatment and/or study from the time of enrollment (start of treatment).
GF	Growth Factor	ID, PNUM, LNUM	Documents any use of growth factor use during the study period (off-protocol event).



Other Study Datasets

Kevs	Description
ID	A record for each protocol exemption received.
	Summary of Serious Adverse Events submitted to the Virahep-
	C Safety Officer. One record per event containing expedited
	review determination criteria as reported by the Safety Officer
ID ONCDATE	and the Clinical Center PI. Can be linked with the adverse
ID, ONSDATE	event record (AE dataset) by onset date.
	One record per patient timepoint with lab results along with upper and lower normal ranges, specific to clinical center and
ID TMPT DSMDATE	date of test.
ID, TWF 1, DSWIDATE	One record per PEG-interferon event (cap opening). Non-
	events (cap openings for refills or accidental openings)
	recorded on the TW are removed. If a patient ever received a
	replacement cap, data was incorporated into the MEMSINT
ID	dataset under the original cap serial number.
	One record per ribavirin event (cap opening). Non-events (cap
	openings for refills or accidental openings) recorded on the TW
	are removed. If a patient ever received a replacement cap,
ID.	data was incorporated into the MEMSRIB dataset under the
ID ID	original cap serial number. Genotype results for screened patients. One record per patient.
	Genotype results for screened patients. One record per patient. Genotype results are by VERSANT™ HCV Genotype Assay
	(Bayer, Tarrytown, NY) completed by BBI/Seracare. Dr. John
	Tavis (virology ancillary) provided results if the genotype
ID, DSMDATE	required sequencing.
	Quantitative and Qualitative test results.* One record per
	patient timepoint. Includes dilution results for quantitative
	testing and 'intent to treat' variable to accommodate one error
ID, DSMDATE	with the originally reported Screen 1 viral level.
	Qualitative test results by patient timepoint for every qualitative
	test performed by BBI/Seracare. Each record contains up to two qualitative results and the associated internal control
	results. Results with negative internal control results are
	included, thus explaining some of the retests. Qualitative tests
	are currently run in duplicate but testing procedures have
	changed throughout the study. There may be more than one
ID, DSMDATE	record per patient timepoint.
	Serum Interferon levels (pg/ml)** for each patient over the
	treatment period: day 0, 1, 2, 3, 7, 14, 28, and treatment week
	8, 12, 24, and 48. Serum interferon levels were also measured
ID TMPT	at Follow-up week 12. Day 3 was not collected after 7/7/2003
ID, TIVIET	due to protocol change. The serum concentrations of 2'-5' OAS** are measured
	(pmol/dL) for each patient during the treatment period: day 0, 1,
	2, 3, 7, 14, 28 and treatment week 8, 12, 24 and 48.
	Comment flags are BLQ = Below the Limit of Quantitation
	AAR = Above the accepted Range; S = Single value. Day 3
ID, TMPT	was not collected after 7/7/2003 due to protocol change.
	ID, ONSDATE ID, TMPT, DSMDATE ID ID ID ID, DSMDATE ID, DSMDATE ID, DSMDATE

^{*} Quantitative test: COBAS Amplicor Hepatitis C Virus Monitor Test, version 2.0 TM assay, (sensitivity 600 IU/ml: Roche Molecular Diagnostics, Alameda, CA)
Qualitative test: Amplicor TM assay (sensitivity 50 IU/ml. Roche)

**OAS and IFN assays were performed by MDS Pharma Services