

Entered to eCRF

Cardiovascular Cell Therapy Research Network TIME Protocol Workbook

FORM NO. CNB005						
Acrostic Identifier:						
Study ID:	۸. /	1				
Date form completed		veical Eva	m (Day of I	nfusion - 3	Day Group)	
Date of Exam: /	/		s outside tii		Reason:	
Date of Exam	_/]	S outside til	ne window		
	Vital S	_			NYHA Class:	
Weight:		pounds				
Temperature:	°F	oral	auricle		□ II	
Respirations:	breath	s/minute			<u></u> III	
Heart rate:	bea	its/minute			□ IV	
Blood Pressure:	/ SBP	 DBP	mmHg (sup	ine)	□ N/A	
Review of Systems:	ОБІ	וטטו				
Have changes occurred	since previou	e vicit?	Yes N	o 🔲 If no,	the table is complete.	
nave changes occurred	Since previou	2 AISIL:	163		the table is complete.	
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>Desc</u>	<u>cribe</u>
Skin						
HEENT						
Lungs						
CV						
Abdomen						
Lymph Nodes						
Musculoskeletal						
Neurological						
Other:						
Questions:						
Has the patient experien	ced any adve	rse events?			Vac 🗆 Na 🗆	
(If yes, complete AE form	n)				Yes ∐ No ∐	
Have there been any cha	•	ications?			Yes No	
(If yes, update medication			4- 4b O I	-1-0		
Was the "Day 3" MRI co (If no, please explain in t			to the Core L	_ab?	Yes 🔲 No 🗌	
			or EACS one	lycic and		
Were five 10 ml venous blood (purple top tubes) for FACS analysis and						
one 10 ml venous blood (green top heparin tube) for plasma cryostorage drawn to ship to the biorepository?						
=	(If no, please explain in Comments)					
Verify patient consented to Biorepository before you draw						
Biorepository bloods.	* * * *					
Was one 3 ml yellow top	tube (anti-co	agulated wit	th acid citrate	e dextrose)		
	for preparation/blinding of the placebo product drawn and sent to the cell Yes No processing lab? (If no, please explain in Comments)					
Comments:	icasc expiain	III COMMINICI)			
Commonto.	Comments:					

Initials _____



FORM NO. CNB005)					
Acrostic Identifier:						
Study ID:						
Date form completed	d: /	/				
		ysical Exa	m (Day of I	nfusion - 7	Day Group)	
Date of Exam: /	1	☐ Visit i	is outside ti	me window	Reason:	
	Vital S	Signs			NYHA Class:	
Weight:		pounds				
Temperature:	°F	oral	auricle			
Respirations:	breath	s/minute				
Heart rate:	bea	nts/minute			□ IV	
Blood Pressure:	/ SBP	 DBP	mmHg (sup	ine)	□ N/A	
Review of Systems:	3=:					
Have changes occurred	since previou	ıs visit?	Yes N	o 🔲 If no,	the table is complete.	
Review of Systems	·	Normal	Abnormal	Not Examined	,	<u>cribe</u>
Skin						
HEENT						
Lungs						
CV						
Abdomen						
Lymph Nodes						
Musculoskeletal						
Neurological						
Other:						
Questions:						
Has the patient experien		erse events?	1		Yes 🗌 No 🗌	
(If yes, complete AE forr		0				
Have there been any cha (If yes, update medication	-	ications?			Yes No	
Was the "Day 3" MRI co		results sent	to the Core I	_ab?	Vac D Na D	
(If no, please explain in	the Comment	s)			Yes No L	
Were five 10 ml venous						
one 10 ml venous blood (green top heparin tube) for plasma cryostorage						
drawn to ship to the biorepository? (If no places symbols in Comments) Yes No						
(if no, please explain in Comments)						
Verify patient consented to Biorepository before you draw Biorepository bloods.						
Was one 3 ml yellow top	tube (anti-co	agulated wit	th acid citrate	e dextrose)		
for preparation/blinding	of the placebo	product dra	awn and sent		Yes 🗌 No 🗌	
processing lab? (If no, p	lease explain	in Commen	ts)			
Comments:						

Entered to eCRF Initials _____



FORM NO. CNB029		
Acrostic Identifier:		
Study ID:		
Date form completed:/	/	
В	one Marrow Aspirat	ion
Procedure Date:		
Procedure Venue:	☐ Patient Room	☐Cath Lab ☐OR
Time initial aspiration start:	:	
Time aspiration complete:	:	
Total amount aspirated:	ml	
Did the patient experience an the procedure? (If yes, complete	_	Yes □No □
Were concomitant medications given? (If yes, add to Medication form) Yes □No □		Yes □ No □
Comments:		
Entered to eCRF	Initials	



FORM NO. CNB031				
Acrostic Identifier:				
Study ID:				
Date form completed:/				
Vital Signs Pre-Cath (Pre-S	tudy Product Infusion)			
Date:/	Time::			
Temperature:	°F 🗌 oral	☐ auricle		
Respirations:	breaths/minute			
Heart rate:	beats/minute			
Blood Pressure:	/	mmHg (sup	ine)	
Study Product Infusion Per				
Procedure Start Date:/_		::		
Stop Date:/_		::		
Was the revascularized vess at the time of cell administration	. NO 1 1	Yes □	If no, pt is ineligible for continuation; fill out an AE and End of Study form	
ST segment changes?	No 🗌	Yes 🗌		
Nitroglycerin given?	No 🗌	Yes 🗌	Amount: mcg (IC)	
Heparin given?	No 🗌	Yes 🗌	Amount: units	
Infusion Catheter Information	:			
Manufacturer:				
Model Name:				
Model Number:				
Diameter:	mm			
Infusion 1 Start Date:/_	_/ Start Time:	<u>:</u>	Volume of infusion 1: ml	
Stop Date:/_	/ Stop Time:	:	☐ Not done	
Infusion 2 Start Date:/_	_/ Start Time:	<u> </u>	Volume of infusion 2: ml	
Stop Date:/_	/ Stop Time:	<u>:</u>	☐ Not done	
Infusion 3 Start Date:/_	_/ Start Time:	:	Volume of infusion 3: ml	
Stop Date:/_	/ Stop Time:	<u>:</u>	☐ Not done	
Infusion 4 Start Date:/_	_/ Start Time:	:	Volume of infusion 4: ml	
Stop Date:/_	/ Stop Time:	:	☐ Not done	
Infusion 5 Start Date:/_	_/ Start Time:	:	Volume of infusion 5: ml	
Stop Date:/_	/ Stop Time:	:	☐ Not done	
Infusion 6 Start Date:/_	_/ Start Time:	:	Volume of infusion 6: ml	
Stop Date:/_	/ Stop Time:	:	☐ Not done	



FORM NO. CNB031					
Acrostic Identifier:					
Study ID:					
Date form completed:/	/				
Vital Signs Post-Cath (Post-	Study Product Infusion				
Date://	Time::				
Temperature:	°F □ oral	auricle			
Respirations:	breaths/minute				
Heart rate:	beats/minute				
Blood Pressure:	/ SBP DBP	mmHg (sup	oine)		
Questions					
1. Was the procedure premat (If yes, complete AE or SAE, a		Yes □	No 🗌		
2. Was the procedure restarted?		Yes 🗌	No 🗌	N/A 🗌	
3. Did the patient experience an adverse event during the procedure? (If yes, complete AE or SAE form)		Yes □	No 🗌		
Were concomitant medications given? (If yes, add to Medication form)		Yes □	No 🗌		
Comments:					
Entered to eCRE	Initials				



FORM #	DESCRIPTION of TIME FORMS	TIME Excel Wkbk tab name
CNB099	Screening/Demographics	Enrollmt
CNB001	Eligibility	Elig
CNB003	Baseline Risk Factors & Other Cardiac Hx	Risk
CNB004	Baseline Non Cardio. Med. Hx	Med Hx
CNB005	Physical Exams	BSL PE/PE
CNB006	Index Event (Revascularization)	PCI
CNB007	Treatment Checklist	Treatment
CNB011	Medication List	Meds
CNB012	Medication Allergies	Meds
CNB021	Labs (Panels)	BSL Labs/D1 Labs
CNB022	Labs (F/U)	Labs M 6,12,24
CNB023	Holter	Holter
CNB024	ECG	ECG
CNB026	Labs (Interim)	Interim Labs
CNB029	Bone Marrow Aspiration	Aspir
CNB031	Study Product Infusion	SPI
CNB041	Adverse Event	AE
CNB042	Serious Adverse Event	SAE
CNB043	Unanticipated Problem	UP
CNB044	Protocol Deviation	Prot Dev
CNB045	Schedule of Procedures	Sched
CNB047	Data Glossary	Glossary
CNB048	Missing Form	Missing
CNB051	End of Study	End



TIME Protocol Work	JOOK
FORM NO. CNB045	
Acrostic Identifier:	
Study ID:	
Schedule of Procedur	res TIME 3-Day Group
Procedures	Time Window
Screening/Baseline	
Screen/Demographics Eligibility (Inclusion/Exclusion criteria) Revascularization/PCI Baseline PE Baseline Lab Panels Baseline Non-Cardiovascular History Baseline Risk Factors	
Baseline Allergies Baseline Medications Baseline EKG Baseline Echo (core) Treatment Checklist	MI - 2 days -/ 4 day
Aspiration/Infusion (SPI) Day of Infusion PE	MI + 3 days +/- 1 day
Biorepository blood draws (if consented) Bone Marrow Aspiration Baseline cMRI (core) Cell Processing Cell Processing - Post Release Study Product Infusion	
Day after Infusion	SPI + 1
Day after Infusion PE Biorepository blood draws (if consented) Day after Infusion Lab Panels Day after Infusion EKG	5.111
1 Month	SPI + 30 days +/- 7 days
PE Labs (F/U) Biorepository blood draws (if consented) EKG Holter	
3 Month	SPI + 90 days +/- 14 days
PE Labs (F/U) Biorepository blood draws (if consented)	
6 Month	SPI + 180 days +/- 30 days
PE Labs (F/U) EKG Biorepository blood draws (if consented) Echo (core) cMRI (core)	



TIME Protocol Workbook

FORM NO. CNB045			
Acrostic Identifier:			
Study ID:			
Schedule of Procedu	res TIME 3-Day Group		
Procedures	Time Window		
12 Month	SPI + 365 days +/- 30 days		
PE Labs (F/U) cMRI*			
24 Month	SPI + 730 days +/- 30 days		
PE Labs (F/U) cMRI* End of Study			

<u>cMRI</u>: all pts. undergo cMRI at Day 3 and pts. Rz to therapy Day 7 undergo repeat cMRI on Day 7 (Day 3 MRI=baseline) <u>Laboratory tests:</u>

<u>Baseline:</u> CBC/diff, lipid panel, renal panel, hepatic panel, CK, CK-MB, Troponin T or I, BNP, hsCRP, pregnancy test for childbearing females (For TIME, all but hsCRP and pregnancy test will be done as part of routine care)

<u>Day 1:</u> CBC/diff, renal panel, hepatic panel, [CK, CK-MB, Troponin T or I one time on the morning following infusion] <u>Mo 1,3:</u> CBC/diff, hepatic panel

Mo 6,12,24: CBC/diff, hepatic panel, & BNP

^{*}echo at 12 & 24 months if cMRI is contraindicated



THINE I TOLOCOL WOLK	
FORM NO. CNB045	
Acrostic Identifier:	
Study ID:	
Schedule of Procedu	res TIME 7-Day Group
Procedures	Time Window
Screening/Baseline	
Screen/Demographics	
Eligibility (Inclusion/Exclusion criteria)	
Revascularization/PCI	
Baseline PE	
Baseline Lab Panels	
Baseline Non-Cardiovascular History	
Baseline Risk Factors	
Baseline Allergies	
Baseline Medications	
Baseline EKG	
Baseline Echo (core)	
Treatment Checklist	
Day 3	MI + 3 days +/- 1 day
Baseline cMRI (core)	lill + 5 days +/- 1 day
Aspiration/Infusion (SPI)	MI + 7 days +/- 1 day
Day of Infusion PE	liii i i dayo ii i day
Biorepository blood draws (if consented)	
Bone Marrow Aspiration	
cMRI	
Cell Processing	
Cell Processing - Post Release	
Study Product Infusion	
Day after Infusion	SPI + 1
Day after Infusion PE	0.111
Biorepository blood draws (if consented)	
Day after Infusion Lab Panels	
Day after Infusion EKG	
1 Month	SPI + 30 days +/- 7 days
PE	
Labs (F/U)	
Biorepository blood draws (if consented)	
EKG	
Holter	
3 Month	SPI + 90 days +/- 14 days
PE	
Labs (F/U)	
Biorepository blood draws (if consented)	
6 Month	SPI + 180 days +/- 30 days
PE	
Labs (F/U)	
EKG	
Biorepository blood draws (if consented)	
Echo (core)	
cMRI (core)	
	l .



TIME Protocol Workbook

FORM NO. CNB045	
Acrostic Identifier:	
Study ID:	
Schedule of Procedu	res TIME 7-Day Group
Procedures	Time Window
12 Month	SPI + 365 days +/- 30 days
PE Labs (F/U) cMRI*	
24 Month	SPI + 730 days +/- 30 days
PE Labs (F/U) cMRI* End of Study	

<u>cMRI</u>: all pts. undergo cMRI at Day 3 and pts. Rz to therapy Day 7 undergo repeat cMRI on Day 7 (Day 3 MRI=baseline) <u>Laboratory tests:</u>

<u>Baseline:</u> CBC/diff, lipid panel, renal panel, hepatic panel, CK, CK-MB, Troponin T or I, BNP, hsCRP, pregnancy test for childbearing females (For TIME, all but hsCRP and pregnancy test will be done as part of routine care)

<u>Day 1:</u> CBC/diff, renal panel, hepatic panel, [CK, CK-MB, Troponin T or I one time on the morning following infusion] <u>Mo 1,3:</u> CBC/diff, hepatic panel

Mo 6,12,24: CBC/diff, hepatic panel, & BNP

Baseline

^{*}echo at 12 & 24 months if cMRI is contraindicated



FORM NO. CNB09	9		
Date source/workboo	Date source/workbook completed://		
Scre	eening / Demograp	hics	
Last Name:			
First Name:			
Middle Initial:			
Consent signed	Biorepository consent signed	Yes No	
Date of Birth			
Sex	$M \square F \square$		
Hispanic	N 🗌 Y 🔲		
Race (choose one) White Black or African			
Asian Native Hawaiian or Other Pacific Islander American Indian or Alaska Native			
Marital Status (cho	ose one):		
Married Living with a par Single/never ma Widowed Divorced Separated	tner		
Entered to eCRF	Initials		



FORM	FORM NO. CNB001				
Acrostic Identifier:					
<u> </u>	Study ID:				
Date so	Date source/workbook completed:// Date of PCI://				
		Eligibility Criteria			
	Note: The use of either DES or BMS for percutaneous revascularization of the infarct-artery is required. The				
Y	N	vessel must be patent at the time cell administration is to be attempted. Inclusion Criteria (Must answer Yes to all questions to be eligible)			
		Patient is at least 21 years of age			
		First MI with successful primary PCI in an artery diameter ≥ 2.5 mm within 24 hours of onset			
		of symptoms			
		No contraindications to undergoing cell therapy procedure within 3-7 days post AMI and PCI			
		Hemodynamic stability as defined as not requiring IABP, inotropic or blood pressure supporting medications			
		Ejection fraction following reperfusion with PCI ≤ 45% as assessed by echocardiography			
		Consent signed. Date signed//			
		Women of childbearing potential willing to use an active form of birth control (If male, check "Y")			
Υ	N	Exclusion Criteria (Must answer No to all questions to be eligible)			
		History of sustained ventricular arrhythmias not related to AMI (evidenced by previous Holter monitoring and/or medication history for sustained ventricular arrhythmias in patient's medical chart)			
		Requires CABG or PCI due to the presence of residual coronary stenosis >70% luminal obstruction in the non-infarct related vessel (Additional PCI of non-culprit vessels may be performed prior to enrollment)			
		History of any malignancy within the past 5 years excluding non-melanoma skin cancer or cervical cancer <i>in-situ</i>			
		History of chronic anemia (hemoglobin (Hgb) < 9.0 mg/dl)			
		History of thrombocytosis (platelets > 500k)			
		Baseline platelet count (prior to revascularization) <120,000 or known history of thrombocytopenia			
		Known history of elevated INR (PT) or PTT			
		Life expectancy less than one year			
		History of untreated alcohol or drug abuse			
		Currently enrolled in another investigational drug or device trial			
		Previous CABG			
		Previous MI with resultant LVEF < 55%			
		History of stroke or TIA within the past 6 months			
		History of severe valvular heart disease (aortic valve area < 1.0 cm ² or > 3+ mitral regurgitation)			



FORM NO. CNB001								
Acrostic Identifier:								
Study ID:								
Eligibility Criteria								
Υ	N	Exclusion Criteria continued (Must answer No to all questions to be eligible)						
	☐ ☐ Pregnant or breast feeding							
		Has a known history of HIV, or has active Hepatitis B or C infection, or active TB						
		Has an active inflammatory or autoimmune disease on chronic immunosuppressive therapy						
		Contraindications to cMRI						
	Previous radiation to the pelvis with WBC and platelet counts below hospital-specific normal values							
	Women of childbearing potential not willing to use an active form of birth control (If male, check "N")							
		Chronic liver disease that might interfere with survival or treatment with cell therapy						
		Chronic renal insufficiency as defined by a creatinine ≥ 2.0 mg/dl or requires chronic dialysis						
	•	tient became ineligible during the screening process; not all data were collected to answer uestion; all questions addressed with the patient have been answered						
	has bee	usion or exclusion criteria exemption, or approval for the most recent protocol amendment, en granted by the CCTRN Medical Monitor or IRB respectively on one or more of the above comment required with a brief explanation; include detail if multiple criteria are involved)						
Comme	Comments:							
PI Signature Date:								
RNC Signature Date:								
Entered to eCRF Initials								



FORM NO. CNB003									
Acrostic Identifier:									
Study ID:									
Date source/workbook completed://									
Baseline Risk Factors									
Diabetes	No		Type I		Type II				
<u>Diabetes Treatment:</u>	0	ral Hyp	ooglycemics						
			Insulin						
			Neither						
Hypertension	No		Yes						
Hypertension Treatment:			1 med						
		2 or	more meds						
		nc	medication						
Hyperlipidemia	No		Yes						
<u>Hyperlipidemia Treatment:</u>		Die	et controlled						
		Dru	g controlled						
			Neither						
Family Hx of MI	No		Yes		Unknown				
Angina	No		Stable		Unstable				
Carotid Disease, asymptomatic	No		Yes						
Hx of TIAs	No		Yes						
Hx of valvular heart disease	No		Yes						
If yes, check all that apply:			mitral						
			aortic						
			pulmonic						
			tricuspid						
Hx of aneurysm	No		Yes						
Hx of Stroke	No		current deficit		completely resolved				
Hx of PVD	No		Yes						
Obese or Hx of obesity	No		Yes						
Smoking	Never		Previous		Current				
	INCVCI		Yr stopped:		_ packs/day:				



FORM NO. CNB003											
Acrostic Identifier:											
Study ID:											
Date source/workbook completed:											
Baseline Risk Factors											
Othe	Other Cardiac History										
Prior to this hospitalization, have you been hospitalized for:											
Congestive Heart Failure	No		Yes		/	1					
Revascularizations (non-CABG)	No		Yes		/	1					
Previous MI	No		Yes		/	1					
Bypass surgery (CABG)	No		Yes		/	1					
Cardiac catheterization	No		Yes		/	1					
Cardiac pacemaker	No		Yes		/	1					
Other coronary interventions	No		Yes								
If yes, please describe other coron	ary interv	ention	s:								
Procedure:					Date mo	st recen	ıt:				
1.					/	1					
2.					/	1					
3.					/	1					
Comments:											
Entered to eCRF Initials											



FORM NO. CNB004											
Acrostic Identifier:											
Study ID:											
Date source/workbook completed://											
Baseline Non-Cardiovascular Medical History Not Describe the abnormality											
System	Not discussed	Unremarkable	Abnormal	Describe the abnormality (obligatory if abnormal)							
Ears, Nose, Throat											
Opthalmic											
Respiratory											
GI											
Renal											
Urogenital											
Neurologic											
Endocrine											
Musculoskeletal											
Skin											
Psychiatric											
Other											
Entered to aCRE	Initials										



	M NO. CNB01	2											
	Acrostic Identifier:												
Study													
Date	source/workbo	ook completed	d:/	Madiaat	ion Allow	wie e							
				Please lis	ion Aller	gies							
Dr	Drug Allergies NKDA Yes												
FOR	FORM NO. CNB011												
Acros	stic Identifier:												
Study	y ID:												
Date	form completed	:	<u>/</u>										
		<u> </u>			dications		Prior to						
	Medicatio	n Class	Medication Name	Dose	Unit	Frequency	Study Start	Start Date	Stop Date	Comments			
1													
2													
3													
4													
5													
6													
7													
8													
9													
10													
11													
12													
13													
14										_			
15										_			
16													
17													
18													

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TIME Protocol Workbook



Medication eCRF drop down lists:

Drug Classes

Allopurinol

Angiotensin converting enzyme inhibitors

Antianginal
Antiarrhythmics
Antibiotics
Anticoagulants

Antiplatelet agents (non-aspirin)

Aspirin

Beta blockers

Calcium channel blockers Cholesterol-lowering agents

Digitalis Diuretics Inotrope Insulin Nitrates

Non-ACE inhibitor arterial vasodilators (e.g. hydralazine)

Non-insulin hormones Oral hypoglycemics Other antihypertensives

Pain medications

Potassium Supplements

Sympathetic blockers

Tranquilizers Vaccines Vasodilators Others <u>Units</u> CAP=capsule g=gram GR=grain

GTT=drop
IU=international units
mg=milligram
mL=milliliter
oz=ounce
PUF=puff
SPY=spray/squirt
SUP=suppository
TAB=tablet
TBS=tablespoon
TSP=teaspoon
U=units
ug=microgram

uL=microliter UNK=unknown

OTH=other (specify)

Frequency
BID=twice daily
ONCE=one dose
per hour
per minute
PRN=as needed
QD=once daily
QID=4 times/day
QOD=everyother day
TID=3 times/day
OTH=other (specify)



FORM NO. CNB024		
Acrostic Identifier:		
Study ID:		
Date source/workbook completed://		
	G - Baseline	
Date of Procedure:// Time:	:	
PR interval: 0 sec QRS interval: 0 se		
☐ ECG NORMAL ☐ ECG NOT NORMAL		
Note: If you select "ECG NORMAL", you are done wit	h this form	
•		
Rhythm: (Choose all that apply)	U ventrieuler demand necembles (VA/I)	
normal sinus rhythm	ventricular demand pacemaker (VVI)	
sinus arrhythmia	atrial pacemaker	
sinus bradycardia (<60 bpm)	dual chamber pacemaker (DDD)	
sinus tachycardia (>100 bpm)	wandering pacemaker	
atrial fibrillation	accelerated idioventricular rhythm	
atrial flutter	atrial premature complexes	
multifocal atrial tachycardia	ventricular premature complexes (PVCs)	
supraventricular tachycardia	ventricular couplets	
iunctional tachycardia	junctional rhythm	
ventricular bigeminy	ventricular fibrillation	
ectopic atrial rhythm		
	bpm (must fill in a & b if this box is checked)	
If ventricular tachycardia, please complete:		
a. Length: complexes b. Aver	-	
If patient is on pacemaker (as indicated above), ch		
☐ 100% paced ☐ intermittently paced	N/A (If 100% paced, do not complete res	
AV Conduction Abnormalities (Choose one):		NONE
AV block, 1st degree		
AV block, 2nd degree Mobitz type 1 (Wenke	ebach)	
AV block, 2nd degree Mobitz type 2		
AV block, 3rd degree		
Abnormalities of P wave: (Choose all that apply)		NONE
Left atrial enlargement	Right atrial enlargement	
Abnormalities of QRS axis (Choose one):		NONE
Left axis deviation(> -30°)	Right axis deviation (> +100°)	
QRS voltage abnormalities: (Choose all that apply)		NONE
Low voltage	Right ventricular hypertrophy	
Left ventricular hypertrophy		



FORM NO. CNB024							
Acrostic Identifier:							
Study ID:							
Date source/workbook completed:	/						
		ECC	G - Baselin	е			
Intraventricular conduction abnormalities: (C	hoo	se a	II that apply)			NONE
Right bundle branch block, complet	:e		Left bur	ndle branch bloo	ck, complete		
Right bundle branch block, incompl	ete		Left bur	ndle branch bloo	ck, incomplete		
Left anterior fascicular block			☐ Nonspe	cific intraventrio	cular conduction	disturbance	:
Left posterior fascicular block							
			For eac	h "Yes" respon	se, check all loca	tions that a	pply:
Are Q waves present?	Υ		N 🗌	Anterior	Lateral	Inferior	
Is ST segment elevation present?	Υ		N 🗌	Anterior	Lateral	Inferior	
Is ST segment depression present?	Υ		N 🗌	Anterior	Lateral	Inferior	
Is T wave inversion present?	Υ		N \square	Anterior	Lateral	Inferior	
Is there evidence of posterior infarction?	Υ		N 🗆	Pathologic R wave V ₁ , V ₂	Abn. ST depression V ₁ , V ₂	Abn. ST elevation V ₁ , V ₂	
Is there evidence of RV infarction (right precordial leads)?	Y		N 🗆	N/A 🗌			
Are there nonspecific ST and/or T wave abnormalities present?	Y		N 🗆				
Comments:							
PI Signature					Date:		
Entered to eCRF Initials							



FORM NO. CNB006								
Acrostic Identifier:								
Study ID:								
Date source/workbook completed://								
Revascularization / PCI								
Date of onset of chest pain:		1	Time::					
Date patient presented at ER:	/	1	Time::					
Date of PCI:		1	Time::					
Calculated ischemic period:	(program	n calculates	s time between or	nset of chest pain	and PCI)			
Calculated door-to-ballon time:	Calculated door-to-ballon time: (program calculates time between presented at ER and PCI)							
TIMI Flow & TMP scores	F	Pre PCI: ₋		Post PCI:				
Artery Location of PCI (Set	ect all that	apply)	Stent Type					
Artery:	<u>No</u>	<u>Yes</u>	<u>None</u>	Drug eluting	Bare metal			
Circumflex								
RCA								
LAD								
Note: If more than one stent is placed in an artery, please state the number of stents per location(s) in the Comments section.								
Comments:								
Entered to eCRE	Initials							



Entered to eCRF Initials _____

Cardiovascular Cell Therapy Research Network TIME Protocol Workbook

FORM NO. CNB005											
Acrostic Identifier:											
Study ID:											
Date source/workbook completed:/											
Physical Exam (Baseline)											
Date of Exam://											
Vital Signs NYHA Class:											
Height:		inches				1					
Weight:		pounds				II					
Temperature:	°F	oral	auricle			Ш					
Respirations:	breath	ns/min				IV					
Heart rate:	bea	ats/min				N/A					
Blood Pressure:	/			ino)		_ % (screening echo)					
Blood Fressure.	SBP	DBP	mmHg (sup	iiie)		evidence of LV n the screening echo?	☐Yes ☐No				
Review of Systems:											
<u>Organs</u>		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>Descri</u>	<u>be (</u> must describe if ans	wer abnormal)				
Skin											
HEENT											
Lungs											
cv											
Abdomen											
Lymph Nodes											
Musculoskeletal											
Neurological											
Other:	Other:										
Questions											
Has the patient experienced any adverse events? (If yes, complete AE form) Yes No											
Was the baseline echo completed to send to the Core Lab? (If no, please enter a reason in Comments) Yes No											
Comments:											

Workbooks Version 5 07/19/2010 Baseline



FORM NO. CNB021									
Acrostic Identifier:									
Study ID:									
Date source/workbook completed://									
Laboratory Tests (Baseline)									
Date and time specimen obta	ained: Da	ate:	// Time::						
CBC with Differential	Result	Unit	Normal Range						
WBC		K/mm ³	4.0-11.0 K/mm ³						
RBC		M/mm ³	4.0-6.0 M/mm ³						
Hgb		gm/dL	12.0-17.5 gm/dL						
Hct		%	33-53%						
MCV		fL	78-100 fL						
Platelets		K/mm ³	135-450 K/mm ³						
WBC Differential									
Neutrophilis		%	36-74%						
Lymphocytes		%	12-45%						
Monocytes		%	0-13%						
Eosinophils		%	0-8%						
Basophils		%	< 3.0%						
Cardiac Enzymes (Either Tropo	onin T or Tro	ponin I shou	ld be completed, NOT both.)						
Troponin T		ng/ml	0.0-10 ng/ml						
Troponin I		ng/ml	0.0-100 ng/ml						
CK		U/L	25-10,000 U/L						
CK-MB		ng/ml	0.0-250 ng/ml						
Renal Panel									
Na+		mmol/L	132-148 mmol/L						
K+		mmol/L	3.3-5.5 mmol/L						
Chloride		mmol/L	95-110 mmol/L						
CO ₂		mmol/L	22-32 mmol/L						
Glucose		mg/dL	65-110 mg/dL						
BUN		mg/dL	5-26 mg/dL						
Creatinine		mg/dL	0.4-1.5 mg/dL						
Albumin		g/dL	3.5-5.0 gm/dL						
Calcium		mg/dL	8.0-10.6 mg/dL						
Inorganic Phosphorus		mg/dL	2.5-4.7 mg/dL						



FORM NO. CNB021								
Acrostic Identifier:								
Study ID:								
Date source/workbook completed://								
Laboratory Tests (Baseline)								
Hepatic Panel	Result	Unit	Normal Range					
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL					
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL					
Total Protein		g/dL	6.0-8.5 g/dL					
Alk Phos		U/L	30-150 U/L					
ALT		U/L	0-50 U/L					
AST		U/L	0-42 U/L					
Lipid Panel			•					
Cholesterol		mg/dL	100-240 mg/dL					
Triglycerides		mg/dL	0-200 mg/dL					
HDL Cholesterol		mg/dL	32-95 mg/dL					
Calculated LDL		mg/dL	60-129 mg/dL					
Chol / HDL Ratio			0-4.5					
Other Tests								
BNP		pg/ml	0-100 pg/ml					
hsCRP		mg/L	0.0-40 mg/L					
Pregnancy Test (women of childbearing age)			Negative (urine)					
☐ Not applicable age or gender			< 5.0 mU/ml (quantitative blood)					
Comments:								
PI Signature								
Entered to eCRF	Initials							



FORM NO. CNB007								
Acrostic Identifier:								
Study ID:								
Treatment Checklist								
Date source/workbook co	Date source/workbook completed: / /							
Please enter a Value from Physical Exam, Baseline			• •	y, Baseline				
Variable	Value		Criteria					
Patient Age		Must be ≥ 21 years old	at consent date					
LVEF		Must be ≤ 45%						
Temperature		Must be < 100.4 °F						
Hemoglobin		Must be ≥ 9.0 mg/dl						
Platelets*		Must be ≥ 120K and ≤ \$	500K BEFORE revaso	cularization*				
Creatinine		Must be < 2.0 mg/dl						
LV thrombus evidence								
Atrial Fibrillation								
If any of the variables above have changed since the Baseline Physical Exam or Baseline Lab Tests, and a more recent exam or test has been done, please enter the updated value, date, and time of the re-check. *Use a platelet count BEFORE revascularization. If the platelet count above was post-revascularization, please enter a pre-revascularization count, test date and time below.								
Variable	Value	Date	Time					
LVEF								
Temperature								
Hemoglobin								
Platelets*								
Creatinine								
A baseline testing exemption has been granted by the CCTRN Medical Monitor for one or more of the above variables, excluding LV thrombus or Afib (comment required with a brief explanation; include detail if multiple variables are involved). Answers to questions below cannot be overridden by checking this box.								



FORM NO. CNB007			
Acrostic Identifier:			
Study ID:			
Treatment Checklist			
Please answer the following questions:			
1. Since the baseline exam and tests, has there been a change in the properties of t	patient's	Yes 🗌	No 🗌
2. Is there any other reason you think this patient should not continue i study? (If yes, please explain in the Comments)	n the	Yes	No 🗌
3. If the patient has an LV thrombus or atrial fibrillation, does either correquire current anticoagulation therapy?	ndition	Yes 🗌	No 🗌
4. Have there been any additional tests that have revealed evidence of LV thrombus that requires anticoagulation therapy?			No 🗆
Comments:			
RNC Signature	Date:		
PI Signature	Date:		
Entered to eCRF Initials			



FORM NO. CNB005						
Acrostic Identifier:						
Study ID:						
Date source/workbo	ok complet	ted:/	/			
			xam (Day a		n)	
Date of Exam: /	1		is outside vindow	Reason:		
☐ Informed consent	was revised	since stud	y start date			
Date patient reconsent	ed: <u>/</u>	1	Consent	version:		
	Vital	Signs			NYHA Class:	
Weight:		pounds				
Temperature:	°F	oral	auricle		П	
Respirations:		ns/minute				
Heart rate:	be	ats/minute			□ IV	
Blood Pressure:	/ SBP	 DBP	mmHg (sup	ine)	□ N/A	
Review of Systems:						
Have changes occurred	since previou	ıs visit?	Yes	No 🗌	(If no, skip to telemetry)	
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>Describe</u>	
Skin						
HEENT						
Lungs						
cv						
Abdomen						
Lymph Nodes						
Musculoskeletal						
Neurological						
Other:						
Telemetry						
If intervention was required, select arrhythmia that required intervention (see list on page 2)		Describe int	ervention:			
If intervention was required, select arrhythmia that required intervention (see list on page 2)			Describe int	ervention:		
If intervention was required, select arrhythmia that required intervention (see list on page 2)			Describe int	ervention:	Workbooks Version 5	



FORM NO. CNB005				
Acrostic Identifier:				
Study ID:				
Date source/workbook completed://				
Physical Exam (Day after Infusion))			
Questions				
Has the patient experienced an adverse event since receiving study product? (If yes, complete AE form)	Yes No			
Have there been any changes to medications since receiving study product? (If yes, update medication form)	Yes No			
Have there been any EKG changes from baseline? (see EKG form)	Yes No No			
If yes, are the changes clinically significant? (see EKG form)	Yes No No			
Was the temperature log given to the patient?	Yes No No			
Were two 10 ml venous blood (purple top tubes) for FACS analysis and one 10 ml venous blood (green top heparin tube) for plasma cryostorage drawn to ship to the biorepository? (If no, please explain in Comments) Verify patient consented to Biorepository before you draw Biorepository bloods.	Yes No 🗆			
Comments:				
<u>Arrhythmias:</u> sinus tachycardia, atrial fibrillation/flutter, accelerated idioventricular rhythm, multifocal atrial tachycardia, supraventricular tachycardia, junctional tachycardia/rhythm, ventricular fibrillation				
Entered to eCRF Initials				



FORM NO. CNB021						
Acrostic Identifier:						
Study ID:						
Laborato	ry Tests (I	Day after l	Infusion)			
Date source/workbook completed:/_	Date source/workbook completed://					
Date and time specimen obtained: D	ate:	/ /	Time::			
CBC with Differential	Result	Unit	Normal Range			
WBC		K/mm ³	4.0-11.0 K/mm ³			
RBC		M/mm ³	4.0-6.0 M/mm ³			
Hgb		gm/dL	12.0-17.5 gm/dL			
Hct		%	33-53%			
MCV		fL	78-100 fL			
Platelets		K/mm ³	135-450 K/mm ³			
WBC Differential						
Neutrophilis		%	36-74%			
Lymphocytes		%	12-45%			
Monocytes		%	0-13%			
Eosinophils		%	0-8%			
Basophils		%	< 3.0%			
Cardiac Enzymes (Either Troponin T or Tro	ponin I sho	uld be comp	eleted, NOT both.)			
Troponin T (Time::)		ng/ml	0.0-10 ng/ml			
Troponin I (Time::)		ng/ml	0.0-100 ng/ml			
CK (Time::)		U/L	25-10,000 U/L			
CK-MB (Time::)		ng/ml	0.0-250 ng/ml			
Renal Panel	Result	Unit	Normal Range			
Na+		mmol/L	132-148 mmol/L			
K+		mmol/L	3.3-5.5 mmol/L			
Chloride		mmol/L	95-110 mmol/L			
CO ₂		mmol/L	22-32 mmol/L			
Glucose		mg/dL	65-110 mg/dL			
BUN		mg/dL	5-26 mg/dL			
Creatinine		mg/dL	0.4-1.5 mg/dL			
Albumin		g/dL	3.5-5.0 gm/dL			
Calcium		mg/dL	8.0-10.6 mg/dL			
Inorganic Phosphorus		mg/dL	2.5-4.7 mg/dL			



FORM NO. CNB021				
Acrostic Identifier:				
Study ID:				
L	aboratory Tests	(Day after	Infusion)	
Hepatic Panel				
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL	
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL	
Total Protein		g/dL	6.0-8.5 g/dL	
Alk Phos		U/L	30-150 U/L	
ALT		U/L	0-50 U/L	
AST		U/L	0-42 U/L	
Comments:				
DI Cignosturo			Doto	
PI Signature			Date:	
Entered to eC	RF Initials _			



FORM NO. CNB024
Acrostic Identifier:
Study ID:
ECG (Day after Infusion)
Date source/workbook completed:/
Date of Procedure: / / Time: :
PR interval: 0 sec QRS interval: 0 sec QT interval: 0 sec HR: bpm
☐ ECG NORMAL ☐ ECG NOT NORMAL
Note: If you select "ECG NORMAL", you are done with this form.
Rhythm: (Choose all that apply)
normal sinus rhythm ventricular demand pacemaker (VVI)
sinus arrhythmia atrial pacemaker
sinus bradycardia (<60 bpm) dual chamber pacemaker (DDD)
sinus tachycardia (>100 bpm) wandering pacemaker
atrial fibrillation accelerated idioventricular rhythm
atrial flutter atrial premature complexes
multifocal atrial tachycardia ventricular premature complexes (PVCs)
supraventricular tachycardia ventricular couplets
☐ junctional tachycardia ☐ junctional rhythm
ventricular bigeminy ventricular fibrillation
ectopic atrial rhythm
ventricular tachycardia (< 30 seconds) > 120 bpm (must fill in a & b if this box is checked)
If ventricular tachycardia, please complete:
a. Length: complexes b. Average Rate: bpm
If patient is on pacemaker (as indicated above), choose level of pacing:
☐ 100% paced ☐ intermittently paced ☐ N/A (If 100% paced, do not complete rest of form
AV Conduction Abnormalities (Choose one):
AV block, 1st degree
AV block, 2nd degree Mobitz type 1 (Wenkebach)
AV block, 2nd degree Mobitz type 2
AV block, 3rd degree
Abnormalities of P wave: (Choose all that apply)
☐ Left atrial enlargement ☐ Right atrial enlargement
Abnormalities of QRS axis (Choose one):
☐ Left axis deviation(> -30°) ☐ Right axis deviation (> +100°)
QRS voltage abnormalities: (Choose all that apply)
☐ Low voltage ☐ Right ventricular hypertrophy
Left ventricular hypertrophy



FORM NO. CNB024							
Acrostic Identifier:							
Study ID:							
	ECG (Day	after Infu	sion)			
Intraventricular conduction abnormalities: (Choose	all	that apply)			[NONE
Right bundle branch block, comple	ete		Left bun	dle branch bloc	k, complete		
Right bundle branch block, incomp	plete		Left bun	dle branch bloc	k, incomplete		
Left anterior fascicular block			Nonspec	cific intraventric	ular conduction	disturbance	
Left posterior fascicular block							
			For each	n "Yes" respons	se, check all loca	ations that a	pply:
Are Q waves present?	Υ [N 🗌	Anterior	Lateral 🗌	Inferior	
Is ST segment elevation present?	Υ [N \square	Anterior	Lateral 🗌	Inferior	
Is ST segment depression present?	Υ [N 🗌	Anterior	Lateral 🗌	Inferior	
Is T wave inversion present?	Υ [N \square	Anterior	Lateral 🗌	Inferior	
Is there evidence of posterior infarction?	Υ [N 🗆	Pathologic R wave V ₁ , V ₂	Abn. ST depression V ₁ , V ₂	Abn. ST elevation V_1, V_2	
Is there evidence of RV infarction (right precordial leads)?	Υ [N 🗆	N/A 🔲			
Are there nonspecific ST and/or T wave abnormalities present?	Υ [N 🗆				
Comments:							
PI Signature				Date:	·		



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Baseline Screening** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1.	CNB099	Screening/Demographics						
2.	CNB001	Eligibility						
3.	CNB003	Baseline Risk Factors & Other Cardiac History						
4.	CNB004	Baseline Non Cardiovascular Medical History						
5.	CNB011	Medication List*	Medication List*					
6.	CNB012	Medication Allergies*						
7.	CNB024	ECG (Baseline)						
8.	CNB006	Index Event (Revascularization)						
9.	CNB005	Baseline Physical Exam						
10	. CNB021	Baseline Laboratory Tests						
11	. CNB007	Treatment Checklist						
*CNB(011 and CNB(012 are on the same page						
Signat	ure		ate					
Printe	d Name							



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Aspiration/Infusion** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1.	CNB005	Day of Infusion Physical Exam		
2.	CNB029	Bone Marrow Aspiration		
3.	CNB031	Study Product Infusion		
Cianatu			Doto	
Signatu	re		Date	
Printed	Name			



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Day after Infusion** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1. CNB005	Day after infusion Physical Exam		
2. CNB021	Day after Infusion Laboratory Tests		
3. CNB024	ECG		
Signature		Date	
Printed Name			



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Month 1 Follow-up Visit** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1.	CNB005	Month 1 Physical Exam		
2.	CNB022	Month 1 Laboratory Tests		
3.	CNB024	ECG		
4.	CNB023	Holter		
Signat	ure		Date	
Printe	d Name			



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Month 3 Follow-up Visit** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1. CNB005 2. CNB022	Month 3 Physical Exam Month 3 Laboratory Tests		
Signature		Date	
Printed Name			



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Month 6 Follow-up Visit** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1. CNB005	•	
2. CNB022	,	
3. CNB024	4 ECG	
Signature		Date
Printed Name		



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Month 12 Follow-up Visit** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

 CNB005 CNB022 	Month 12 Physical Exam Month 12 Laboratory Tests		
Signature		Date	
Printed Name			



TIME Protocol Workbook

FORM COMPLETION ATTESTATION

All of the following forms associated with **Month 24 Follow-up Visit** have been completed and entered as electronic case report forms in the CCTRN web application, or a missing form has been noted.

1. CNB005	Month 24 Physical Exam	
2. CNB022	Month 24 Laboratory Tests	
Signature		Date
Printed Name		



FORM NO. CNB005							
Acrostic Identifier:							
Study ID:							
Date source/workbo	ok complet	ed:/_	/				
		Physica	I Exam (Mo	onth 1)			
Date of Exam: / / Reason: time window							
☐ Informed consent	was revised	since stud	y start date				
Date patient reconsent	ed: <u>/</u>	1	Consent	version:			
	Vital	Signs			NYHA Class:		
Weight:		pounds			I		
Temperature:	°F	oral	auricle				
Respirations:	breath	ns/minute					
Heart rate:	bea	ats/minute			□ IV		
5	/				□ N/A		
Blood Pressure:	SBP	DBP	mmHg (sup	ine)			
Review of Systems:							
Have changes occurred	since previou	ıs visit?	Yes	No 🗌	(If no, table is complete))	
Review of Systems		<u>Normal</u>	Abnormal	Not Examined	<u>Describe</u>		
Skin							
HEENT							
Lungs							
CV							
Abdomen							
Lymph Nodes							
Musculoskeletal							
Neurological							
Other:							



TIME Protocol Workbook

FORM NO. CNB005		
Acrostic Identifier:		
Study ID:		
Date source/workbook completed://		
Physical Exam (Month 1)		
Questions		
Has the patient experienced a new adverse event since the last follow-up visit? (If yes, complete AE form)	Yes No [
Have there been any changes to medications since the last follow-up visit? (If yes, update medication form)	Yes No	
Have there been any ECG changes from baseline? (see ECG form)	Yes No [
If yes, are the changes clinically significant? (see ECG form)	Yes No [
Were there any significant findings on the Holter report? (see Holter form)	Yes No [
Were two 10 ml venous blood (purple top tubes) for FACS analysis and one 10 ml venous blood (green top heparin tube) for plasma cryostorage drawn to ship to the biorepository? (If no, please explain in Comments) Verify patient consented to Biorepository before you draw Biorepository bloods.	Yes □ No □	
Comments:		
Entered to eCRF Initials		



FORM NO. CNB022							
Acrostic Identifier:							
Study ID:							
Date source/workbook completed:/							
Laboratory Tests (Month 1)							
Date and time specimen obtained: Date:/ Time: :							
CBC with Differential	Result	Unit	Normal Range				
WBC		K/mm ³	4.0-11.0 K/mm ³				
RBC		M/mm ³	4.0-6.0 M/mm ³				
Hgb		gm/dL	12.0-17.5 gm/dL				
Hct		%	33-53%				
MCV		fL	78-100 fL				
Platelets		K/mm ³	135-450 K/mm ³				
WBC Differential							
Neutrophilis		%	36-74%				
Lymphocytes		%	12-45%				
Monocytes		%	0-13%				
Eosinophils		%	0-8%				
Basophils		%	< 3.0%				
Hepatic Panel							
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL				
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL				
Total Protein		g/dL	6.0-8.5 g/dL				
Alk Phos		U/L	30-150 U/L				
ALT		U/L	0-50 U/L				
AST		U/L	0-42 U/L				
Comments:			•				
PI Signature			Date:				
Entered to eCRF	Initials						



FORM NO. CNB024	
Acrostic Identifier:	
Study ID:	
Date source/workbook completed:/	
ECG (Month 1)	
Date of Procedure: / / Time: :	
PR interval: 0 sec QRS interval: 0 sec QT interval: 0 sec HR: bpm	
☐ ECG NORMAL ☐ ECG NOT NORMAL	
Note: If you select "ECG NORMAL", you are done with this form.	
Rhythm: (Choose all that apply)	
normal sinus rhythm ventricular demand pacemaker (VVI)	
sinus arrhythmia atrial pacemaker	
sinus bradycardia (<60 bpm) dual chamber pacemaker (DDD)	
sinus tachycardia (>100 bpm) wandering pacemaker	
atrial fibrillation accelerated idioventricular rhythm	
atrial flutter atrial premature complexes	
multifocal atrial tachycardia ventricular premature complexes (PVCs)	
supraventricular tachycardia ventricular couplets	
junctional tachycardia junctional rhythm	
ventricular bigeminy ventricular fibrillation	
ectopic atrial rhythm	
ventricular tachycardia (< 30 seconds) > 120 bpm (must fill in a & b if this box is checked)	
If ventricular tachycardia, please complete:	
a. Length: complexes b. Average Rate: bpm	
If patient is on pacemaker (as indicated above), choose level of pacing:	
☐ 100% paced ☐ intermittently paced ☐ N/A (If 100% paced, do not complete rest	t of form)
AV Conduction Abnormalities (Choose one):	NONE
AV block, 1st degree	
AV block, 2nd degree Mobitz type 1 (Wenkebach)	
AV block, 2nd degree Mobitz type 2	
AV block, 3rd degree	
Abnormalities of P wave: (Choose all that apply)	NONE
Left atrial enlargement Right atrial enlargement	
Abnormalities of QRS axis (Choose one):	NONE
☐ Left axis deviation(> -30°) ☐ Right axis deviation (> +100°)	
QRS voltage abnormalities: (Choose all that apply)	NONE
☐ Low voltage ☐ Right ventricular hypertrophy	
Left ventricular hypertrophy	



FORM NO. CNB024							
Acrostic Identifier:							
Study ID:							
Date source/workbook completed:/		/_					
		EC	G (Month 1)			
Intraventricular conduction abnormalities: (Ch	100	se a	I that apply				NONE
Right bundle branch block, complete	Э		Left bur	idle branch bloc	k, complete		
Right bundle branch block, incomple	ete		Left bur	dle branch bloc	k, incomplete		
Left anterior fascicular block			☐ Nonspe	cific intraventric	ular conduction o	listurbance	!
Left posterior fascicular block							
			For eac	h "Yes" respons	se, check all loca	tions that a	pply:
Are Q waves present?	Υ		N 🔲	Anterior	Lateral 🗌	Inferior	
Is ST segment elevation present?	Υ		N \square	Anterior	Lateral 🗌	Inferior	
Is ST segment depression present?	Υ		N 🔲	Anterior	Lateral 🗌	Inferior	
Is T wave inversion present?	Υ		N 🔲	Anterior	Lateral 🗌	Inferior	
Is there evidence of posterior infarction?	Υ		N 🗆	Pathologic R wave V ₁ , V ₂	Abn. ST depression V ₁ , V ₂	Abn. ST elevation V ₁ , V ₂	
Is there evidence of RV infarction (right precordial leads)?	Υ		N 🗆	N/A 🔲			
Are there nonspecific ST and/or T wave abnormalities present?	Υ		N 🔲				
Comments:							
PI Signature				Date	:		



Entered to eCRF

Initials _____

Cardiovascular Cell Therapy Research Network

TIME Protocol Workbook

FORM NO. CNB023						
Acrostic Identifier:						
Study ID:						
Date source/workbook completed://						
Holter Data Form <i>(Month 1)</i>						
Date procedure started:/Predominant Rhythm: (mutually exclusive						
Total recording time:::	Sinus Rhythm					
General:	☐ Paced Rhythm ☐ Ectopic Atrial Rhythm					
Total beats/QRS Complexes: beats	Atrial Flutter / Fibrillation					
Paced beats: beats	Heart Rates:					
Pauses/Longest RR Interval (> 2 secs):	Minimum:beats/min. @:					
Longest pause was seconds @:	Average:beats/min.					
Total number of pauses:	Maximum:beats/min. @:					
Ventricular Arrhythmia Summary:	Supraventricular Arrhythmia Summary:					
Single/PVC: beats	Single/PAC: beats					
Couplets:	Couplets:					
Total number of NSVT Runs (≥ 3 beats)	Total number of SVT Runs					
Number of beats in longest NSVT run	Number of beats in longest SVT run					
Total number of sustained ventricular tachycardia runs	Intermittent Atrial Fibrillation / Atrial Flutter:					
(≥ 30 secs)	☐ Yes ☐ No					
	If yes, total no. of episodes					
	If yes, min.secs (duration of longest episode)					
AV Block: (Choose all that apply)						
☐ Transient AV block, 2nd degree-Mobitz type 1 (Wenket	pach)					
total no. of episodes do	uration of longest episode (secs)					
☐ Transient AV block, 2nd degree-Mobitz type 2 ☐ N/A						
total no. of episodes d	uration of longest episode (secs)					
☐ Transient AV block, 3rd degree ☐ N/A						
total no. of episodes d	uration of longest episode (secs)					
Comments:						
PI Signature	Date:					



FORM NO. CNB005							
Acrostic Identifier:							
Study ID:							
Date source form co	mpleted: _	/	/				
		Physica	al Exam (Mo	,			
Date of Exam: / / Visit is outside time window Reason:							
☐ Informed consent	was revised	since stud	y start date				
Date patient reconsent	ed: <u>/</u>	1	Consent	version:			
Vital Signs NYHA Class:							
Weight:		pounds			I		
Temperature:	°F	oral	auricle				
Respirations:		ns/minute			III		
Heart rate:	bea	ats/minute					
Blood Pressure: / mmHg (supine) SBP DBP					∐ N/A		
Review of Systems:							
Have changes occurred	since previou	ıs visit?	Yes 🗌	No 🗌	(If no, table is complete)	1	
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>Describe</u>		
Skin							
HEENT							
Lungs							
CV							
Abdomen							
Lymph Nodes							
Musculoskeletal							
Neurological							
Other:							



FORM NO. CNB005		
Acrostic Identifier:		
Study ID:		
Date source form completed://		
Physical Exam (Month 3)		
Questions		
Has the patient experienced a new adverse event since the last follow-up visit? (If yes, complete AE form)	Yes 🗌 No [
Have there been any changes to medications since the last follow-up visit? (If yes, update medication form)	Yes No [
Were two 10 ml venous blood (purple top tubes) for FACS analysis and one 10 ml venous blood (green top heparin tube) for plasma cryostorage drawn to ship to the biorepository? (If no, please explain in Comments) Verify patient consented to Biorepository before you draw Biorepository bloods.	Yes □ No [
Comments:		
Entered to eCRF Initials		



FORM NO. CNB022			
Acrostic Identifier:			
Study ID:			
Date form completed: ////			
Labo	oratory Te	ests (Mont	h 3)
Date and time specimen obtained: D	ate:	/ /	Time: :
CBC with Differential	Result	Unit	Normal Range
WBC		K/mm ³	4.0-11.0 K/mm ³
RBC		M/mm ³	4.0-6.0 M/mm ³
Hgb		gm/dL	12.0-17.5 gm/dL
Hct		%	33-53%
MCV		fL	78-100 fL
Platelets		K/mm ³	135-450 K/mm ³
WBC Differential			
Neutrophilis		%	36-74%
Lymphocytes		%	12-45%
Monocytes		%	0-13%
Eosinophils		%	0-8%
Basophils		%	< 3.0%
Hepatic Panel			
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL
Total Protein		g/dL	6.0-8.5 g/dL
Alk Phos		U/L	30-150 U/L
ALT		U/L	0-50 U/L
AST		U/L	0-42 U/L
Comments:			
PI Signature			Date:
Entered to eCRF	Initials		



FORM NO. CNB005									
Acrostic Identifier:									
Study ID:									
Date form completed	d:								
Physical Exam (Month 6)									
Date of Exam:/		U Visit i	s outside tir	me window	Reason:				
☐ Informed consent was revised since study start date									
Date patient reconsente	ed: <u>/</u>	1	Consent	version:					
	Vital	Signs			NYHA Class:				
Weight:		pounds							
Temperature:	°F	oral	auricle		<u> </u>				
Respirations:	breath	ns/minute			III				
Heart rate:	bea	ats/minute			□IV				
Blood Pressure:	/ SBP	 DBP	mmHg (sup	ine)	□ N/A				
Review of Systems:									
Have changes occurred	since previou	ıs visit?	Yes No	☐ If no, th	ne table is complete.				
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>De</u>	<u>escribe</u>			
Skin									
HEENT									
Lungs									
CV									
Abdomen									
Lymph Nodes									
Musculoskeletal									
Neurological									
Other:									



FORM NO. CNB005			
Acrostic Identifier:			
Study ID:			
Date form completed:			
	Physical Exam (Month 6)		
Questions			
Has the patient experienced an adver product? (If yes, complete AE form)	se event since receiving study	Yes No	
Have there been any changes to med product? (If yes, update medication for	- ,	Yes No	
Have there been any ECG changes fr (see ECG form)	Yes No		
If yes, are the changes clinically sig	nificant? (see ECG form)	Yes \square No	
Was the Echo completed to send to tl (If no, please enter a reason in Comm		Yes No	
Was the MRI completed to send to th (If no, please enter a reason in Comm		Yes 🗌 No	MRI contraindicated
Were two 10 ml venous blood (purple one 10 ml venous blood (green top he drawn to ship to the biorepository? (If no, please explain in Comments) Verify patient consented to Biorepository bloods.	eparin tube) for plasma cryosto-rage	Yes □ No	
Comments:			



FORM NO. CNB022				
Acrostic Identifier:				
Study ID:				
Date source/workbook completed:				
La	aboratory Te	ests (Mont	th 6)	
Date and time specimen obtained: Date:/ Time::				
CBC with Differential	Result	Unit	Normal Range	
WBC		K/mm ³	4.0-11.0 K/mm ³	
RBC		M/mm ³	4.0-6.0 M/mm ³	
Hgb		gm/dL	12.0-17.5 gm/dL	
Hct		%	33-53%	
MCV		fL	78-100 fL	
Platelets		K/mm ³	135-450 K/mm ³	
WBC Differential				
Neutrophilis		%	36-74%	
Lymphocytes		%	12-45%	
Monocytes		%	0-13%	
Eosinophils		%	0-8%	
Basophils		%	< 3.0%	
Hepatic Panel				
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL	
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL	
Total Protein		g/dL	6.0-8.5 g/dL	
Alk Phos		U/L	30-150 U/L	
ALT		U/L	0-50 U/L	
AST		U/L	0-42 U/L	
Other Tests				
BNP		pg/ml	0-100 pg/ml	
Comments:				
PI Signature			Date:	
Entered to eCRF	Initials			



FORM NO. CNB024	
Acrostic Identifier:	
Study ID:	
Date source/workbook completed://	
ECG (Month 6)	
Date of Procedure: / / Time::	
PR interval: 0 sec QRS interval: 0 sec QT interval: 0 sec	sec HR: bpm
☐ ECG NORMAL ☐ ECG NOT NORMAL	
Note: If you select "ECG NORMAL", you are done with this form.	
Rhythm: (Choose all that apply)	
normal sinus rhythm ventricular demand	pacemaker (VVI)
sinus arrhythmia atrial pacemaker	
sinus bradycardia (<60 bpm) dual chamber pacer	maker (DDD)
sinus tachycardia (>100 bpm) wandering pacemak	` ,
atrial fibrillation accelerated idiovent	
atrial flutter atrial premature con	•
multifocal atrial tachycardia ventricular prematur	re complexes (PVCs)
supraventricular tachycardia ventricular couplets	
☐ junctional tachycardia ☐ junctional rhythm	
ventricular bigeminy ventricular fibrillation	n
ectopic atrial rhythm	
ventricular tachycardia (< 30 seconds) > 120 bpm (must fill in a & b if	f this box is checked)
If ventricular tachycardia, please complete:	
a. Length: complexes b. Average Rate: bpm	1
If patient is on pacemaker (as indicated above), choose level of pacing:	
☐ 100% paced ☐ intermittently paced ☐ N/A (If 100% paced	paced, do not complete rest of form)
AV Conduction Abnormalities (Choose one):	□NONE
AV block, 1st degree	
AV block, 2nd degree Mobitz type 1 (Wenkebach)	
AV block, 2nd degree Mobitz type 2	
AV block, 3rd degree	
Abnormalities of P wave: (Choose all that apply)	□NONE
Left atrial enlargement Right atrial enlargen	ment
Abnormalities of QRS axis (Choose one):	□NONE
☐ Left axis deviation(> -30°) ☐ Right axis deviation	·
QRS voltage abnormalities: (Choose all that apply)	NONE
☐ Low voltage ☐ Right ventricular hyp	pertrophy
Left ventricular hypertrophy	



Acrostic Identifier: Study ID: Date source/workbook completed:/	FORM NO. CNB024					
Date source/workbook completed:/	Acrostic Identifier:					
Intraventricular conduction abnormalities: (Choose all that apply)	Study ID:					
Intraventricular conduction abnormalities: (Choose all that apply) Right bundle branch block, complete Right bundle branch block, incomplete Left bundle branch block, incomplete Left anterior fascicular block Left anterior fascicular block For each "Yes" response, check all locations that apply: Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Is there evidence of posterior infarction? Are there nonspecific ST and/or T wave abnormalities present? NONE Left bundle branch block, complete Left bundle branch block, incomplete Left bundle branch block, complete Left bundle branch block, incomplete Left bundle branch block.	Date source/workbook completed:/_	/_				
Right bundle branch block, complete Right bundle branch block, incomplete Left anterior fascicular block Left posterior fascicular block For each "Yes" response, check all locations that apply: Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is T wave inversion present? Is there evidence of Posterior infarction? Right bundle branch block, complete Left bundle branch block, incomplete Left bundle branch block, incomplete Left bundle branch block, complete Left bundle branch block, incomplete Nonspecific intraventricular conduction disturbance For each "Yes" response, check all locations that apply: Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Pathologic Abn. ST depression V1, V2 V1, V2 V1, V2 V1, V2 Is there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?		EC	G (Month 6	;)		
Right bundle branch block, incomplete Left anterior fascicular block Left posterior fascicular block For each "Yes" response, check all locations that apply: Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Is there evidence of Posterior infarction? Are evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Intraventricular conduction abnormalities: (Choo	se a	ll that apply)		NONE
Left anterior fascicular block □ Nonspecific intraventricular conduction disturbance Left posterior fascicular block For each "Yes" response, check all locations that apply: Are Q waves present? Y □ N □ Anterior □ Lateral □ Inferior □ Is ST segment elevation present? Y □ N □ Anterior □ Lateral □ Inferior □ Is ST segment depression present? Y □ N □ Anterior □ Lateral □ Inferior □ Is T wave inversion present? Y □ N □ Anterior □ Lateral □ Inferior □ Is there evidence of posterior infarction? Y □ N □ Pathologic Abn. ST Abn. ST R wave depression elevation V1, V2 □ V2, V2 □ V2, V2 □ V2, V3, V4 Is there evidence of RV infarction (right precordial leads)? Y □ N □ N/A □ Are there nonspecific ST and/or T wave abnormalities present? Y □ N □	Right bundle branch block, complete		Left bur	ndle branch bloc	k, complete	
Left posterior fascicular block For each "Yes" response, check all locations that apply: Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Pathologic Abn. ST Abn. ST R wave depression elevation V ₁ , V ₂ V ₁ , V ₂ Is there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Right bundle branch block, incomplete		Left bur	ndle branch bloc	k, incomplete	
For each "Yes" response, check all locations that apply: Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Abn. ST R wave depression elevation V ₁ , V ₂ V ₂ , V ₃ , V ₄ Is there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Left anterior fascicular block		☐ Nonspe	cific intraventric	ular conduction o	disturbance
Are Q waves present? Y N Anterior Lateral Inferior Is ST segment elevation present? Y N Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Inferior Is T wave inversion present? Y N Anterior Anterior Detailed Inferior Inferior Inferior Is T wave inversion present? N R wave depression V1, V2 Department of the precordial leads)? Are there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Left posterior fascicular block					
Is ST segment elevation present? Y N Anterior Lateral Inferior Is ST segment depression present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Pathologic Abn. ST Abn. ST R wave depression elevation Y, V,			For eac	h "Yes" respons	e, check all loca	tions that apply:
Is ST segment depression present? Y N Anterior Lateral Inferior Is T wave inversion present? Y N Anterior Lateral Inferior Pathologic Abn. ST Abn. ST R wave depression V ₁ , V ₂ N depression V ₁ , V ₂ N N N N N N N N N N N N N N N N N N N	Are Q waves present?		N 🗌	Anterior	Lateral 🗌	Inferior
Is T wave inversion present? Y N Anterior Lateral Inferior Pathologic R wave depression elevation V ₁ , V ₂ N N N/A Is there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Is ST segment elevation present? Y		N 🗌	Anterior	Lateral	Inferior
Is there evidence of posterior infarction? Y N Pathologic R wave depression V ₁ , V ₂ N depression V ₁ , V ₂ N N N/A N/A N/A N/A N/A N/A N/A N/A N/A	Is ST segment depression present?		N 🗌	Anterior	Lateral	Inferior
Is there evidence of posterior infarction? Y \square N \square R wave depression V_1, V_2 \square ls there evidence of RV infarction (right precordial leads)? Are there nonspecific ST and/or T wave abnormalities present?	Is T wave inversion present? Y		N 🗌	Anterior	Lateral 🗌	Inferior
precordial leads)? Are there nonspecific ST and/or T wave abnormalities present? Y N N	Is there evidence of posterior infarction?		N 🗆	R wave	depression	elevation
abnormalities present?			N 🗆	N/A 🔲		
Comments:			N 🗆			
	Comments:					
PI Signature Date:	_			Date:		



FORM NO. CNB005						
Acrostic Identifier:						
Study ID:						
Date form completed	d:/	/				
		Physical	Exam (Mo	nth 12)		
Date of Exam:/ _/						
☐ Informed consent	was revised	since stud	y start date			
Date patient reconsent	ed: <u>/</u>	1	Consent	version:		
	Vital S	Signs			NYHA Class:	
Weight:		pounds			□ I	
Temperature:	°F	□oral	auricle			
Respirations:	breath	ns/minute				
Heart rate:	bea	ats/minute			□ IV	
Blood Pressure:	/ SBP	— — — DBP	mmHg (sup	ine)	□ N/A	
Review of Systems:						
Have changes occurred	since previou	ıs visit?	Yes	No 🗌	(If no, table is complete)	
Review of Systems		<u>Normal</u>	Abnormal	Not Examined	<u>Describe</u>	
Skin						
HEENT						
Lungs						
cv						
Abdomen						
Lymph Nodes						
Musculoskeletal						
Neurological						
Other:						



FORM NO. CNB005	
Acrostic Identifier:	
Study ID:	
Date form completed:/	
Physical Exam (Month 12)	
Questions	
Has the patient experienced a new adverse event since the last follow-up visit? (If yes, complete AE form)	Yes No
Have there been any changes to medications since the last follow-up visit? (If yes, update Medication form)	Yes No
Was the safety MRI completed? (If no, enter a reason in Comments)	Yes No
Was the safety Echo completed? (The safety Echo is only required if the MRI is contraindicated) (If both safety MRI and safety Echo not done then a comment is required)	Yes No
Comments:	
Entered to eCRE	



FORM NO. CNB022			
Acrostic Identifier:			
Study ID:			
Date form completed://			
La	boratory Te	sts (Monti	h 12)
Date and time specimen obtained: Date:/ Time: :			
CBC with Differential	Result	Unit	Normal Range
WBC		K/mm ³	4.0-11.0 K/mm ³
RBC		M/mm ³	4.0-6.0 M/mm ³
Hgb		gm/dL	12.0-17.5 gm/dL
Hct		%	33-53%
MCV		fL	78-100 fL
Platelets		K/mm ³	135-450 K/mm ³
WBC Differential			
Neutrophilis		%	36-74%
Lymphocytes		%	12-45%
Monocytes		%	0-13%
Eosinophils		%	0-8%
Basophils		%	< 3.0%
Hepatic Panel			
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL
Total Protein		g/dL	6.0-8.5 g/dL
Alk Phos		U/L	30-150 U/L
ALT		U/L	0-50 U/L
AST		U/L	0-42 U/L
Other Tests			
BNP		pg/ml	0-100 pg/ml
Comments:			
PI Signature			Date:
Entered to eCRF	Initials		



FORM NO. CNB005							
Acrostic Identifier:							
Study ID:							
Date form completed	d:/	/					
		Physical	Exam (Mo				
Date of Exam:/	Date of Exam: / / / Sist is outside time window						
☐ Informed consent	was revised	since stud	y start date				
Date patient reconsent	ed:/		Consent	version:			
	Vital S	3igns			NYHA Class:		
Weight:		pounds					
Temperature:	°F	□oral	auricle				
Respirations:	breath	ns/minute					
Heart rate:	bea	ats/minute			□ IV		
Blood Pressure:	/ SBP	— — — DBP	mmHg (supi	ine)	□ N/A		
Review of Systems:	CEI						
Have changes occurred	since previou	s visit?	Yes	No 🗌	(If no, table is complete)		
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	Not Examined	<u>Describe</u>		
Skin							
HEENT							
Lungs							
CV							
Abdomen							
Lymph Nodes	Lymph Nodes						
Musculoskeletal							
Neurological							
Other:							



FORM NO. CNB005		
Acrostic Identifier:		
Study ID:		
Date form completed: / /		
Physical Exam (Month 24)		
Questions		
Has the patient experienced a new adverse event since the last follow-up visit? (If yes, complete AE form)	Yes No]
Have there been any changes to medications since the last follow-up vist? (If yes, update Medication form)	Yes No]
Was the safety MRI completed? (If no, enter a reason in Comments)	Yes No]
Was the safety Echo completed? (The safety Echo is only required if the MRI is contraindicated) (If both safety MRI and safety Echo not done then a comment is required)	Yes No]
Comments:		
Entered to aCRE Initials		



FORM NO. CNB022				
Acrostic Identifier:				
Study ID:				
Date form completed:/ _/				
La	boratory Te	sts (Monti	h 24)	
Date and time specimen obtained: Date:// Time: :				
CBC with Differential	Result	Unit	Normal Range	
WBC		K/mm ³	4.0-11.0 K/mm ³	
RBC		M/mm ³	4.0-6.0 M/mm ³	
Hgb		gm/dL	12.0-17.5 gm/dL	
Hct		%	33-53%	
MCV		fL	78-100 fL	
Platelets		K/mm ³	135-450 K/mm ³	
WBC Differential				
Neutrophilis		%	36-74%	
Lymphocytes		%	12-45%	
Monocytes		%	0-13%	
Eosinophils		%	0-8%	
Basophils		%	< 3.0%	
Hepatic Panel				
Bilirubin-Total		mg/dL	0.0-1.5 mg/dL	
Bilirubin-Direct		mg/dL	0.0-0.4 mg/dL	
Total Protein		g/dL	6.0-8.5 g/dL	
Alk Phos		U/L	30-150 U/L	
ALT		U/L	0-50 U/L	
AST		U/L	0-42 U/L	
Other Tests				
BNP		pg/ml	0-100 pg/ml	
Comments:				
PI Signature			Date:	
Entered to eCRF	Initials			



FORM NO. CNB	005							
Acrostic Identifie								
Study ID:								
Date form compl	eted:	/ /						
	Physical Exam (Interim)							
Date of Exam:/		☐ Visit is	outside time	window	Reason:			
☐Informed cons	sent was rev	vised since s	tudy start da	te	Date patient reconsent Consent version:	ted: / /		
	\	/ital Signs			NYHA Class:			
Weight:		pounds			□ I			
Temperature:	°F	oral	auricle					
Respirations:	breat	hs/minute						
Heart rate:	be	ats/minute			□ IV			
Blood Pressure:	/ SBP	 DBP	mmHg (supir	ne)	□ N/A LVEF:%			
Review of Systems	:							
Have changes occur	rred since pr	evious visit?	Yes I		able is complete.			
Review of Systems		<u>Normal</u>	<u>Abnormal</u>	<u>Not</u> Examined	<u>Desc</u>	<u>rribe</u>		
Skin								
HEENT								
Lungs								
CV								
Abdomen								
Lymph Nodes								
Musculoskeletal								
Neurological								
Other:								
Questions								
Has the patient experienced a new adverse event since the last follow-up visit? (If yes, complete AE form)								
Have there been any changes to medications since the last follow-up visit? (If yes, update medication form)								
Comments:								
Entered to eCRF Initials								



FORM NO. CNB026						
Acrostic Identifier:						
Study ID:						
Date form completed:/						
	Laboratory Te	ests (Interi	m)			
Reason for Interim Lab:						
Date and time specimen obtained	ed: Date:	/ /	Time:::			
CBC with Differential	Result	Unit	Normal Range			
WBC		K/mm ³	4.0-11.0 K/mm ³			
RBC		M/mm ³	4.0-6.0 M/mm ³			
Hgb		gm/dL	12.0-17.5 gm/dL			
Hct		%	33-53%			
MCV		fL	78-100 fL			
Platelets		K/mm ³	135-450 K/mm ³			
WBC Differential						
Neutrophilis		%	36-74%			
Lymphocytes		%	12-45%			
Monocytes		%	0-13%			
Eosinophils		%	0-8%			
Basophils		%	< 3.0%			
Cardiac Enzymes (Either Tropon	nin T or Troponin	I should be o	completed, NOT both.)			
Troponin T		ng/ml	0.0-10 ng/ml			
Troponin I		ng/ml	0.0-100 ng/ml			
СК		U/L	25-10,000 U/L			
CK-MB		ng/ml	0.0-250 ng/ml			
Renal Panel						
Na+		mmol/L	132-148 mmol/L			
K+		mmol/L	3.3-5.5 mmol/L			
Chloride		mmol/L	95-110 mmol/L			
CO ₂		mmol/L	22-32 mmol/L			
Glucose		mg/dL	65-110 mg/dL			
BUN		mg/dL	5-26 mg/dL			
Creatinine		mg/dL	0.4-1.5 mg/dL			
Albumin		g/dL	3.5-5.0 gm/dL			
Calcium		mg/dL	8.0-10.6 mg/dL			
Inorganic Phosphorus		mg/dL	2.5-4.7 mg/dL			



Bilirubin-Total mg/dL 0.0-1.5 m Bilirubin-Direct mg/dL 0.0-0.4 m Total Protein g/dL 6.0-8.5 g/ Alk Phos U/L 30-150 U/ ALT U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel mg/dL 100-240 r Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg/ Negative Pregnancy Test (women of childbearing age) Negative	FORM NO. CNB026					
Date form completed:/	Acrostic Identifier:					
Laboratory Tests (Interim) Hepatic Panel Result Unit Normal R Bilirubin-Total mg/dL 0.0-1.5 m Bilirubin-Direct mg/dL 0.0-0.4 m Total Protein g/dL 6.0-8.5 g/ Alk Phos U/L 30-150 U/L AST U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Not applicable age or gender Negative < 5.0 mU/L	Study ID:					
Hepatic Panel Result Unit Normal R Bilirubin-Total mg/dL 0.0-1.5 m Bilirubin-Direct mg/dL 0.0-0.4 m Total Protein g/dL 6.0-8.5 g/ Alk Phos U/L 30-150 U/ ALT U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel mg/dL 100-240 r Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ml hsCRP mg/L 0.0-40 mg/ml Negative Pregnancy Test (women of childbearing age) Negative < 5.0 mU/ml	Date form completed:/	/				
Bilirubin-Total mg/dL 0.0-1.5 m	Laboratory Tests (Interim)					
Bilirubin-Direct mg/dL 0.0-0.4 m Total Protein g/dL 6.0-8.5 g/ Alk Phos U/L 30-150 U/ ALT U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel mg/dL 100-240 m Cholesterol mg/dL 100-240 m Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ml hsCRP mg/L 0.0-40 mg/ml Negative Pregnancy Test (women of childbearing age) Negative < 5.0 mU/ml	Hepatic Panel F	Result	Unit	Normal Range		
Total Protein g/dL 6.0-8.5 g/ Alk Phos U/L 30-150 U/A ALT U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Negative < 5.0 mU/A	3ilirubin-Total		mg/dL	0.0-1.5 mg/dL		
Alk Phos	3ilirubin-Direct		mg/dL	0.0-0.4 mg/dL		
ALT U/L 0-50 U/L AST U/L 0-42 U/L Lipid Panel Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/c	Total Protein		g/dL	6.0-8.5 g/dL		
AST U/L 0-42 U/L Lipid Panel Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/ceptor constructions of the construction of the con	Alk Phos		U/L	30-150 U/L		
Lipid Panel Cholesterol mg/dL 100-240 r Triglycerides mg/dL 0-200 mg HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/ Solution 100-240 rg Negative < 5.0 mU/ Negative 100-240 rg	ALT		U/L	0-50 U/L		
Cholesterolmg/dL100-240 rTriglyceridesmg/dL0-200 mgHDL Cholesterolmg/dL32-95 mgCalculated LDLmg/dL60-129 mChol / HDL Ratio0-4.5Other TestsBNPpg/ml0-100 pg/mlhsCRPmg/L0.0-40 mg/mg/LPregnancy Test (women of childbearing age)Negative□ Not applicable age or gender< 5.0 mU/mg/mg/mg/s	AST		U/L	0-42 U/L		
Triglycerides mg/dL 0-200 mg/dL 32-95 mg/dL 32-95 mg/dL 32-95 mg/dL 60-129 m mg/dL 60-129 m O-4.5 Other Tests BNP pg/ml 0-100 pg/msCRP mg/L 0.0-40 mg/dL 0.0-40	_ipid Panel					
HDL Cholesterol mg/dL 32-95 mg Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg Pregnancy Test (women of childbearing age) Not applicable age or gender some significant or construction of the construction o	Cholesterol		mg/dL	100-240 mg/dL		
Calculated LDL mg/dL 60-129 m Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg/ Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/	Friglycerides		mg/dL	0-200 mg/dL		
Chol / HDL Ratio 0-4.5 Other Tests BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg/ Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/	HDL Cholesterol		mg/dL	32-95 mg/dL		
Other Tests BNP	Calculated LDL		mg/dL	60-129 mg/dL		
BNP pg/ml 0-100 pg/ hsCRP mg/L 0.0-40 mg/ Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/	Chol / HDL Ratio			0-4.5		
hsCRP mg/L 0.0-40 mg/Pregnancy Test (women of childbearing age) Not applicable age or gender < 5.0 mU/Pregnancy Test (women of childbearing age)	Other Tests					
Pregnancy Test (women of childbearing age) ☐ Not applicable age or gender Negative < 5.0 mU/	3NP		pg/ml	0-100 pg/ml		
childbearing age) ☐ Not applicable age or gender Negative < 5.0 mU/	nsCRP		mg/L	0.0-40 mg/L		
results ago of genee.	• ,			Negative (urine)		
Comments:	☐ Not applicable age or gender			< 5.0 mU/ml (quantitative blood)		
	Comments:		•			
PI Signature Date:						



TIME Protocol Workbook

FORM NO. CNB	041																	
Acrostic Identifie	r:																	
Study ID:																		
					Adv	/ers	e E	vent	Log									
Date of this Report	::://																	
Outcome Status	Serious	Expectedness	Severity			Study	Onship to // Study Outcome Attributed to AE oduct:					Study Status						
1=Resolved (must have an end date) 2=Ongoing 3=Resulted in SAE (must complete SAE form)	1=Not Serious 2=Serious (must complete SAE form)	1=Expected 2=Unexpected	1=Mild 1=Definite 2=Moderate 2=Probable 3=Severe 3=Possible 4=Life threatening 4=Unlikely or permanently 5=Unrelated 5		1=Resolved, no treatment, no sequelae 2=Resolved, no treatment, with sequelae 3=Resolved with treatment, no sequelae 4=Resolved with treatment and sequelae 5=Still present, no treatment 6=Still present, being treated				1=Continuing in Study 2=Withdrawn									
Descript	ion of Event (Diaວູ	gnosis)	_	Start /	Date /	·)	_	End /	l Date	_)	Outcome Status	Serious	Serious Expectedness Severity Relationship to Study/Study Product to AE			Study Status	Narrative added* (progress note)	
1.																		
2.																		
3.																		
4.																		
5.																		
* Narrative should in	clude the following	: detailed descripti	ion of	f event	, prob	olem,	and	l/or pro	oduct use	е	rror, and	l relevar	nt tests/l	aborato	ry data,	includin	g dates	•
PI Signature						_	Dat	e:										
RNC Signature_								Date:										
Entered to eCRF	Initials																Wor	kbooks Ver



TIME Protocol Workbook

FORM NO. CNB	042										
Acrostic Identifie	Acrostic Identifier:										
Study ID:	Study ID:										
	Serious Adverse Event Log										
Pate of this Report://											
Outcome Status	Expectedness	Severity	Relationship to Study / Study Product:	Outcome Attributed to SAE					Study Status		
1=Resolved (must have an end date) 2=Ongoing	1=Expected 2=Unexpected (may need to fill out Unanticipated Problem (UP) form)	1=Mild 2=Moderate 3=Severe 4=Life threatening or permanently disabling 5=Fatal	1=Definite 2=Probable 3=Possible 4=Unlikely 5=Unrelated	1=Death: (/) 2=Life Threatening 3=Requires or Prolongs Inpatient Hospitalization 4=Persistent or Significant Disability or Incapacity 5=Congenital Anomaly/Birth Defect 6=Other Serious (Important Medical Events)						ing in Study wn	
Description of Event (Diagnosis)			Start Date	End Date (<u>/</u> /_)	Outcome Status	Expectedness	Severity	Relationship to Study/Study Product	Outcome Attributed to SAE	Study Status	Narrative added* (progress note)
1.											
2.											
3.											
4.											
5.	5.										
* Narrative should include the following: detailed description of event, problem, and/or product use error, and relevant tests/laboratory data, including dates											
PI Signature			Date:								
RNC Signature_											
•			Date			_			W	orkbook	s Versio
Entered to eCRF	Entered to eCRF Initials										



FORM I	NO. CNB0	43				
Is this	unanticipa	ated problem specific to an individual subject?	☐ Yes	□No		
Acrostic	c Identifier:	(fill in if answer to above is "Yes")				
Study II	D: <i>(fill in if</i>	answer to above is "Yes")				
Site: (fi	ill in if ansv	ver to above is "No")				
(Note: If	the UP doe	es not apply to an individual subject, the Acrostic Identifier and Si	tudy ID remai	n blank)		
Date for	rm comple	ted://				
		Unanticipated Problem (UP) Report				
		Any problem or event which in the opinion of the local researcher related to the research procedures.	was unantici	oated, serious		
These sh	nould be rep	orted to the IRB within 10 working days.				
Date of	the Event:					
Date the	e site study	team had knowledge of the Event://				
This Eve	ent meets t	the criteria for an unanticipated problem because:				
		Unanticipated: The event is unexpected in terms of nature, se	everity or freq	uency given the		
	1	research procedures described in the protocol, consent, etc. or the population being studied.		, ,		
	2	Related: The event is related or possibly related to participation reasonable possibility that the incident, experience, event, or our caused by the procedures involved in research.				
	3	Serious: The event placed subjects or others at greater risk (in psychological, economic, or social harm) that was previously kn resulted in harm to the subject or others.	•			
Note: Th	he event <u>m</u>	<u>ust meet all</u> of the above criteria to be considered an unanti	icipated prob	olem.		
Describ	e the type					
	to recur.	or unintentional change to the IRB-approved protocol that resulte		·		
	to the risks	in the literature, safety monitoring report, or other findings indicator potential benefits of the research.				
	Complaint or research st	of a participant that indicates an unanticipated risk or which canr aff.	not be resolve	ed by the		
	A breach in computer).	n confidentiality that may involve risk to that individual or others (e.g. comprom	ised/stolen		
	Incarceration of a member of the research staff.					
	Any other event that, in the opinion of the PI, constitutes an unanticipated risk.					
Description of the unanticipated problem:						
	a plan to pro	event the problem from reoccurring in the future (indicate if proto the event):	ocol or conser	nt modifications		
			,	Vorkbooks Versl on		



FORM N	IO. CNB0	44					
Acrostic	Identifier						
Study ID	:						
Date for	m comple	ted://					
		Proto	ocol Deviation/Violation Report				
Date of th	e Event: _	/ /	Event has not yet occurred (exemption request)				
Date the s	site study t	eam had knowledge	of the Event:/				
This Even	This Event meets the criteria for a protocol deviation/violation because:						
	The event resulted in an accidental or unintentional change to the IRB approved protocol and procedures without prior sponsor approval.						
	2	The event affected the	e participant's rights, safety, or welfare, or the integrity of the resultant data.				
Note: The	event mus	st meet at least one o	f the above criteria to be considered a protocol deviation/violation.				
Describe	the proto	ocol deviation/violat	tion:				
Explain v	Explain why or how the deviation/violation occurred:						
Indicate	Indicate the outcome (PI's assessment of the outcome, comments, or determinations):						
Describe	Describe what action you have taken to prevent recurrence:						
PI Signa	ture		Date:				
RNC Sig	RNC Signature Date:						
Entered to eCRF Initials							
CCTRN Exemption/Waiver Documentation (DCC only)							
CCTRN Medical Officer or Designee Review:							
	Action	Taken:	☐ Granted ☐ Not Granted				
Wai	ver Ackn	owledgement:	☐ Received / Acknowledged				
DCC Sig	nature		Date:				



FORM	/I NO. CNB048						
Acros	Acrostic Identifier:						
Study	Study ID:						
	Missing Form						
	Form Missing:	Reason/Comment:	Date of this Entry:				
	BSL Risk Factors & Other Cardiac Hx						
	BSL Non Cardio. Med. Hx						
	BSL - Physical Exam						
	BSL - ECG						
	BSL - Labs						
	Medication allergies						
	Medication list						
	Index Event (Revascularization)						
	Bone Marrow Aspiration						
	Study Product Infusion						
	Day of Infusion - Phys. Exam						
	Day after Infusion - Phys. Exam						
	Day after Infusion - ECG						
	Day after Infusion - Labs						
	Mo 1 - Physical Exam						
	Mo 1 - Labs (F/U)						
	Mo 1 - ECG						
	Holter						
	Mo 3 - Physical Exam						
	Mo 3 - Labs (F/U)						
	Mo 6 - Physical Exam						
	Mo 6 - Labs (F/U)						
	Mo 6 - ECG						
	Mo 12 - Physical Exam						
	Mo 12 - Labs (F/U)						
	Mo 24 - Physical Exam						
	Mo 24 - Labs (F/U)						
	End of Study						



Entered to eCRF

Initials _____

FORM NO. CNB051							
Acrostic Identifier:							
Study ID:							
Date form completed:	/ /						
	End of S	Study					
Date of final follow-up	o study visit://						
Reason for discharge	Reason for discharge from the study:						
☐ Com	☐ Completed study Date of Discharge from Study://						
☐ With	ndrawn Da	te of Discharge from Study://					
☐ Lost	t to follow-up Da	te of Discharge from Study://					
☐ Scre	een Failure Da	te of Discharge from Study:/					
If "Withdrawn", pleas	e check the primary reaso	n for withdrawal:					
Reasons that requi	ire follow-up:						
☐ Serie	ous Adverse Event (until res	solved) Event Number:					
☐ Preg	gnancy (1 year post birth)	Event Number:					
☐ Othe	er	Describe:					
Reasons that DO N	NOT require follow-up:						
☐ Dea	th	Event Number:					
☐ Adve	erse Event	Event Number:					
☐ With	ndrawal of consent						
☐ Prot	ocol Deviation/Violation						
□ Inve	stigator Discretion	Describe:					
☐ Spo	nsor Discretion	Describe:					
☐ Othe	er	Describe:					
Please verify the follo	wing tasks are complete:						
☐ All Informed Con	sents forms are properly sig	ned/dated and available					
☐ Hard copy workb	ooks are signed, dated and	present in the CCTRN source document					
patient binder; w	patient binder; workbooks may be grouped by a visit with one signature per visit.						
☐ All source docum	☐ All source document data have been entered into the electronic CRF database						
☐ All electronic CRFs have been submitted to the DCC							
I have reviewed all case report forms for this patient and found them to be in complete agreement with the source documents.							
	arise from the DCC data rev norized staff will supply appro	riew (due to missing, unclear, or incorrect opriate corrections.					
PI Signature		Date:					
-							