	Non S	erious Adv	erse Drug	Reactions	s Report		
Start Date:2016-09-	-02 End Date:2016-09				•		
Study :EMR200136_583	Investigator :NA	vestigator :NA		SiteNo:007			
Subject No :007-0010	Subject Initials :	DOB :03/25/1989	Sex:Male	Race:Caucasian	Height:179(cm)	Weight:63(kg)	
First administration	tration date of batch :		Batch number :				
Study Drug	Start Date		Dose	Change in Dose	se		
Visit 2 (Month 3)	06/08/2015		9				
Visit 5 (Month12)/Early Termination							
Visit 3 (Month 6)	06/08/2015		9				
Visit 1/ Baseline (Day 1)	06/08/2015		9				
Visit 4 (Month 9)	06/08/2015		9				
Adverse Event	Start Date	End Date	Time related to study treatment	Causality to study drug	Severity		
elevated liver enzymes	09/04/2015			Suspected	Moderate		
Causality Factors		Action Taken with Study Treatment	Other action taken	Outcome	AE Special Interest	AE dose limiting toxicity	
None(Othervalue:)			Concomitant medication	Ongoing			
Event description:		L		1	· L		
Subject received co	ncomitant medication	s:Yes					
Name of medication	Start Date	Ongoing	End Date	Dose	Unit	Frequency	
LIV52	09/04/2015		01/26/2016	6	tb	TID	
Acetaminophen	06/08/2015		06/08/2015	500	mg	PRN	
Solu-Medrol	01/27/2016		01/29/2016	1000	mg	QD	
LIV 52	01/27/2016		02/22/2016	3	tb	TID	
Controloc	01/27/2016		02/03/2016	20	mg	QD	
Rivotril 0.5mg	01/27/2016	Yes		500	mcg	BID	
Milgamma	01/30/2016		02/08/2016	3	tb	TID	
Does the subject ha	ave any relevant past	or present medical cond	litions:No	1	1	1	
Condition				Start Date	Related to study condition	Ongoing	