

BSRBR Event of Special Interest (ESI) Report MI / ACUTE CORONARY SYNDROME

SRBR ritish Society for Rheumatology	ety for Patient Initials:		Gender: Date of Birth: NHS Number:	
ogics Registers matoid Arthritis	Event Date:	Biologic/ bi	osimilar at time of event:	
		Product Batch Number:		
Event Details				
Rise in cardiac markers? YES NO DON'T KNOW				
Trop T/ Trop I Level: (Highest level recorded)				
Did the patient have ischaemic symptoms?				
ECG findings → Were there any ischaemic changes YES NO DON'T KNOW				
→ Were there any new Q waves YES NO DON'T KNOW				
Was the patient thrombolysed?				
Did they receive primary angioplasty on the same day as the event?				
YES NO DON'T KNOW				
Did they have any other cardiac intervention? YES NO DON'T KNOW				
If yes, please specify what & when:				
Do you believe there is a possibility that this adverse event was related to the biologic/biosimilar				
drug used to treat RA? Yes No Unknown				
If Yes please confirm which drug:				
What was the outcome of the event? Resolved Not Resolved Resolved with sequelae Fatal				
Resolved Not Resolved Resolved with sequelae Fatal				
Form completed Return ESI/s to: BSRBR-RA. The University of Manchester,				
By: Unit 4 Rutherford House, 40 Pencroft Way, Manchester Science Park On:/ Manchester, M15 6SZ. You can also fax these to: 0161 2751640				