

BSRBR-RA Event of Special Interest (ESI) CVA

The University of	Manchester
BSR	BR
The British Society for Rheumatology Biologics Registers	
Rheumatoid Arthritis	

Study ID: HRN: Patient Initials:		Gender: Date of Birth: NHS Number:
Event Date:	Biologic/biosimilar at time of event: Product Batch Number:	

		Product Batch Number:
Event Details		
Was the stroke	haemorrhagic	YES NO DON'T KNOW
True the en ene	Or ischaemic	YES NO DON'T KNOW
	Or ischaeniic	TES NO DONTRIOW
Was the patien	t thrombolised?	YES NO DON'T KNOW
Does the patient have atrial fibrillation? YES NO DON'T KNOW		
Or paroxysmal atrial fibrillation? YES NO DON'T KNOW		
Was a CT/MRI done? YES NO DON'T KNOW (If yes, please attach report)		
Did signs/symptoms fully resolve? YES NO DON'T KNOW		
If so, did they resolve within: 24 hours 1 week More than one week		
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 Unknown		
If Yes please confirm which drug:		
What was the outcome of the event? Resolved Not Resolved Resolved with sequelae Fatal		
Form completed By:		Return ESI/s to: BSRBR-RA. The University of Manchester, 4 Rutherford House, 40 Pencroft Way, Manchester Science Park

Manchester, M15 6SZ. You can also fax these to: 0161 2751640