Click here to access the guidelines	/NICE algorithm				
	N&WICS -	TA195 - Infliximab - Rheum	natoid Arthritis		
Before providing patient identifiable parent/legal guardian/carer) has give CSU for processing this funding req	en appropriate explic	cit consent for sensitive person	nal information on this form		
If there is more than one NICE-approabout the advantages and disadvant patient is likely to adhere to treatme and price per dose) unless an order	ages of the treatment. The most appro	ents available. This has taken priate, least expensive, will be	into consideration therapeut	tic need and whethe	er or not the
Patient			Practice		
NHS No:	Trust:		Name:		
Patient Hospital No:	Consultant Making Request:		Practice Postcode:		
Patient's Initials and DoB:			Practice Code:		
Notification Email Address:		(@NHS.net account ONLY)	Contact name & number:		
	Provider:				
Start date of requested treatment:	Supplier:	☐ Private ☐ NHS ☐ Homecare ☐ Hospital	Sub-Type:	/A 🔽	
For support regarding IFRs, please For support regarding the criteria lis			et		
Please indicate whether patier	nt meets the follow	ving NICE criteria:		Please tick	
Rituximab in combination with severe active rheumatoid arthritis disease-modifying anti-rheumatic inhibitor. Treatment with rituximab	who have had an in drugs (DMARDs), i	nadequate response to, or are ncluding at least one tumour i	intolerant of, other necrosis factor (TNF)	□Yes □No	
Treatment with rituximab in column adequate response following initial retreatment with a dosing interval in disease activity score (DAS28)	ation of therapy and of at least 6 months	if an adequate response is mass. An adequate response is de	aintained following	☐Yes ☐No	
3. Adalimumab, etanercept, inflix recommended as treatment option inadequate response to, or have who cannot receive rituximab the is withdrawn because of an adver	ns only for adults wi an intolerance of, ot rapy because they h	th severe active rheumatoid a her DMARDs, including at lea	rthritis who have had an st one TNF inhibitor, and	☐Yes ☐No	
Adalimumab monotherapy and with severe active rheumatoid arthother DMARDs, including at least have a contraindication to methot	nritis who have had one TNF inhibitor,	an inadequate response to, or and who cannot receive rituxing	have an intolerance of, mab therapy because they	☐Yes ☐No	
5. Treatment with adalimumab, e adequate response (as defined in with assessment of DAS28, at lea maintained.	1.2) 6 months after	r initiation of therapy. Treatme	nt should be monitored,	☐Yes ☐No	
6. When using DAS28, healthcar disabilities, communication difficu assessment and make any adjus	ılties, or disease ch	aracteristics that could advers		☐Yes ☐No	
7. A team experienced in the diagnosis and treatment of rheumatoid arthritis and working under the supervision of a rheumatologist should initiate, supervise and assess response to treatment with rituximab,					

adalimumab, etanercept, infliximab or abatacept.	
I confirm that the patient meets the criteria for treatment	
Name of person completing:	Contact Details:
Designation of person completing:	Date:
Trust Authorising Pharmacist	
Name:	
Date:	