Click here to access the guidelines/NICE algorithm

Click here to access the guidelines	/NICE algorithm				
	N&WICS - TA195 - Ad	alimumab - Rheumatoid	Arthritis		
Before providing patient identifiable parent/legal guardian/carer) has give CSU for processing this funding req	n appropriate explicit consent	for sensitive personal infor	mation on this form		
If there is more than one NICE-approabout the advantages and disadvant patient is likely to adhere to treatme	ages of the treatments availab	le. This has taken into con	sideration therapeu	tic need and wheth	er or not the
and price per dose) unless an order			· (talling into accou		ooio, accago
Patient NHS No:	Trust:		Practice Name:		
Patient Hospital No:	Consultant Making Request:		Practice Postcode:		
Patient's Initials and DoB:			Practice Code:		
Notification Email Address:	(@NHS.ne	t account ONLY)	Contact name & number:		
Start date of requested treatment:	0	e □NHS care □Hospital	Sub-Type:	/A 🔽	
commissioning statement. Any requisite For support regarding IFRs, please	contact: norfolkicd@nhs.net		3 1 1 1	,	
Please indicate whether patier	t meets the following NICE	criteria:		Please tick	
Rituximab in combination with severe active rheumatoid arthritis disease-modifying anti-rheumatic inhibitor. Treatment with rituximab	who have had an inadequate durings (DMARDs), including at	esponse to, or are intolerar	nt of, other factor (TNF)	□Yes □No	
Treatment with rituximab in column adequate response following initiar retreatment with a dosing interval in disease activity score (DAS28)	tion of therapy and if an adeq of at least 6 months. An adeq	uate response is maintaine	d following	□Yes □No	
3. Adalimumab, etanercept, inflixi recommended as treatment option inadequate response to, or have who cannot receive rituximab the is withdrawn because of an adver	ns only for adults with severe a an intolerance of, other DMAR apy because they have a con	active rheumatoid arthritis w Ds, including at least one T	ho have had an NF 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 an inadequone TNF inhibitor, and who c	late response to, or have all annot receive rituximab the	n intolerance of, rapy 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 initiation of	of therapy. Treatment shoul	d be monitored,	☐Yes ☐No	
When using DAS28, healthcar disabilities, communication difficu assessment and make any adjus	lties, or disease characteristic	s that could adversely affe		□Yes □No	
7. A team experienced in the diag supervision of a rheumatologist s				☐Yes ☐No	

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:	