Click here to access the guidelines/NICE algorithm

	N&WICS -	TA195 - Etanercept - Rheumat	oid Arthritis		
Before providing patient identifiable parent/legal guardian/carer) has gir CSU for processing this funding re	ven appropriate explic	cit consent for sensitive personal	information on this form		
If there is more than one NICE-app about the advantages and disadva patient is likely to adhere to treatm	ntages of the treatme	ents available. This has taken into	consideration therapeu	tic need and wheth	er or not the
and price per dose) unless an orde	er of preference is sta	ated in the TAs.			
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.net account ONLY)	Contact name & number:		
Start date of requested treatment:	Provider: Supplier:	☐ Private ☐ NHS ☐ Homecare ☐ Hospital	Sub-Type:	/A 🔽	
By completing this form, you confined commissioning statement. Any requirements appeared by the support regarding IFRs, please For support regarding the criterial.	uests which fall outsi	ide of this use will require an indi ⊉nhs.net			
Please indicate whether patie	ent meets the follow	wing NICE criteria:		Please tick	
severe active rheumatoid arthriti disease-modifying anti-rheumati	s who have had an ir c drugs (DMARDs), i	commended as an option for the standequate response to, or are intrincluding at least one tumour nects more frequently than every 6 m	olerant of, other crosis factor (TNF)	□Yes □No	
adequate response following init	tiation of therapy and all of at least 6 month	notrexate should be continued on if an adequate response is main s. An adequate response is defir ore.	tained following	☐Yes ☐No	
recommended as treatment opti inadequate response to, or have	ons only for adults wi an intolerance of, ot erapy because they l	t, each in combination with methith severe active rheumatoid arthither DMARDs, including at least of the account air account of the contraindication to rituxim	ritis who have had an one TNF inhibitor, and	□Yes □No	
with severe active rheumatoid at other DMARDs, including at lea	thritis who have had st one TNF inhibitor,	nerapy are recommended as trea an inadequate response to, or ha and who cannot receive rituxima thotrexate is withdrawn because	ave an intolerance of, b therapy because they	☐Yes ☐No	
adequate response (as defined	in 1.2) 6 months after	o and abatacept should be conting r initiation of therapy. Treatment and continued only if an adequa	should be monitored,	□Yes □No	
	culties, or disease ch	ould take into account any physic aracteristics that could adversely er appropriate.		□Yes □No	
· ·	•	nt of rheumatoid arthritis and wor	-	☐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:	