## Click here to access the guidelines/NICE algorithm

	N&WICS - TA829 - Upadacitinib - Active ankylosin	ng spondylitis				
parent/legal guardian/carer) has give	providing patient identifiable data on this form, please confirm that the patient (or in the case of a minor or vulnerable adult with the egal guardian/carer) has given appropriate explicit consent for sensitive personal information on this form to be passed to the CCG and/or processing this funding request and validating subsequent invoices. Consent given:					
about the advantages and disadvant patient is likely to adhere to treatme	oved treatment available, a discussion between the responsages of the treatments available. This has taken into connt. The most appropriate, least expensive, will be chose of preference is stated in the TAs.	nsideration therapeu	tic need and wheth	er or not the		
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: ☐ Private ☐ NHS  Supplier: ☐ Homecare ☐ Hospital	Sub-Type:	/A 🔽			
For support regarding IFRs, please	ests which fall outside of this use will require an individual contact: norfolkicd@nhs.net ted below, please contact: norfolknontariff@nhs.net	ai runding request (i	rk).			
Please indicate whether patier	nt meets the following NICE criteria:		Please tick			
Upadacitinib is recommended well enough with conventional the	as an option for treating active ankylosing spondylitis the trapy in adults, only if:	at is not controlled				
tumour necrosis factor (TNF)-alg and	oha inhibitors are not suitable or do not control the condi-	tion well enough	□Yes □No			
the company provides upadaciti	nib according to the commercial arrangement.					
	onsider upadacitinib to be one of a range of suitable treat noose the least expensive treatment, taking into accound commercial arrangements.		☐Yes ☐No			
Assess response to upadacitin evidence of response, defined as:	ib after 16 weeks of treatment. Continue treatment only	if there is clear				
a reduction in the Bath Ankylos treatment value or by 2 or more u	ing Spondylitis Disease Activity Index (BASDAI) score to nits and	50% of the pre-	☐Yes ☐No			
a reduction in the spinal pain vis	sual analogue scale (VAS) by 2 cm or more.					
	l, sensory or learning disabilities, or communication diffi Al and spinal pain VAS and make any adjustments nee		□Yes □No			
	ent should normally be started with the least expensive dand any additional administration costs).	Irug (considering				
	drug is considered the most cost-effective treatment for atment options recommended by NICE may be lower in opticable.		☐Yes ☐No			
	reatment is available and has not been tried by the patie oribing has been included in the patients' medical record					

auditing.				
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. J				