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3				
	N&WICS - TA481 - Basiliximab - Kidney transpla	ant in adults		
parent/legal guardian/carer) has give	data on this form, please confirm that the patient (or in the patient explicit consent for sensitive personal info uest and validating subsequent invoices. Consent given:	rmation on this form		
about the advantages and disadvant patient is likely to adhere to treatme	oved treatment available, a discussion between the respi ages of the treatments available. This has taken into coint. The most appropriate, least expensive, will be chose of preference is stated in the TAs.	nsideration therapeu	tic need and wheth	ner 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 individu contact: norfolkicd@nhs.net sed below, please contact: norfolknontariff@nhs.net	arrunding request (i	rk).	
Please indicate whether patier	nt meets the following NICE criteria:		Please tick	
	t of an immunosuppressive regimen that includes a calc to prevent organ rejection in adults having a kidney trar		☐Yes ☐No	
an initial option to prevent organ r started with the least expensive p form if the least expensive produc as a result of a disability or they a reasons). Tacrolimus granules for	when used as part of an immunosuppressive regimen, is ejection in adults having a kidney transplant. Treatment roduct.[3] However, treatment can be started with an altot is not suitable (for example, if the person is not able to re unable to have a particular ingredient because of alle oral suspension (Modigraf) should be used only if the cotat agreed with the Commercial Medicines Unit.	should normally be ernative dosage o swallow capsules rgy or religious	□Yes □No	
3. Mycophenolate mofetil, when used as part of an immunosuppressive regimen, is recommended as an initial option to prevent organ rejection in adults having a kidney transplant. Treatment should normally be started with the least expensive product. However, treatment can be started with an alternative dosage form if the least expensive product is not suitable (for example, if the person is not able to swallow capsules as a result of a disability or they are unable to have a particular ingredient because of allergy or religious reasons).[1],[2]			☐Yes ☐No	
	mmunoglobulin, prolonged-release tacrolimus, mycophe ept are not recommended as initial treatments to preven		☐Yes ☐No	
appraisal as options for preventing technologies recommended in se and a corticosteroid (for example,	make recommendations on any of the technologies consigning organ rejection in adults who are, or become, unable to ctions 1.1 to 1.3 or standard triple therapy with ciclospobecause of treatment failure, contraindications, or intolelcineurin inhibitors, or thrombotic microangiopathy). This	have the rin, azathioprine erance such as		
are unable to continue having the their graft or	eir initial therapy and need to switch to another therapy	during the life of	☐ Yes ☐ No	

• have a second or subsequent transplant, having previously for treatments or standard treatments are clinically unsuitable for econtraindications or intolerance.		
6. As per NICE guidance, treatment should normally be started the dose required, price per dose and any additional administral Please confirm that the choice of drug is considered the most of patient, noting that alternative treatment options recommended use of biosimilar medications if applicable.	cost-effective treatment for this individual by NICE may be lower in cost, including the	□Yes □No
Where an alternative lower cost treatment is available and has not been tried by the patient, please confirm that the clinical rationale for prescribing has been included in the patients' medical record for the purpose of auditing. 7. The product is being used as described by local commissioning position.		
, ,	ing position.	∐Yes ∐No
I confirm that the patient meets the criteria for treatment Name of person completing:	Contact Details:	
Designation of person completing:	Date:	
Designation of person completing: Trust Authorising Pharmacist	Date:	