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	N&WICS - TA755 - Ris	sdiplam - Spinal muscula	ar atrophy		
Before providing patient identifiable of parent/legal guardian/carer) has give CSU for processing this funding requ	en appropriate explicit consent	t for sensitive personal info	rmation 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 treatments availal nt. The most appropriate, leas	ble. This has taken into co st expensive, will be chose	nsideration therapeu	tic need and wheth	er or not the
Patient	or preference is stated in the	17.3.	Practice		
NHS No:	Trust:		Name:		
Patient Hospital No:	Consultant Making Request:		Practice Postcode:		
Patient's Initials and DoB:			Practice Code:		
Notification Email (@NHS.net		et account ONLY)	Contact name & number:		
Start date	Provider:	e □NHS			
of requested treatment:	0	care Hospital	Sub-Type:	/A 🔽	
By completing this form, you confirm commissioning statement. Any requi-	ests which fall outside of this		t described below as		al
For support regarding the criteria list	ted below, please contact; nor	rfolknontariff@nhs.net			
For support regarding the criteria listed below, please contact: norfolknontariff@nhs.net Please indicate whether patient meets the following NICE criteria:					
Risdiplam is recommended as an option for treating 5q spinal muscular atrophy (SMA) in people 2 months and older with a clinical diagnosis of SMA types 1, 2 or 3 or with pre-symptomatic SMA and 1 to 4 SMN2 copies. It is recommended only if the conditions of the managed access agreement are followed.					
2. As per NICE guidance, treatment should normally be started with the least expensive drug (considering the dose required, price per dose and any additional administration costs).					
Please confirm that the choice of drug is considered the most cost-effective treatment for this individual patient, noting that alternative treatment options recommended by NICE may be lower in cost, including the use of biosimilar medications if applicable.					
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.					
3. The product is being used as described by local commissioning position.					
I confirm that the patient meets	s the criteria for treatment				
Name of person completing:		Contact Details:			
Designation of person completing:		Date:			
Trust Authorising Pharmacist					
Name:					