Click here to access the guideline	s/NICE algorithm				
	N&WICS - TA217 -	Memantine - Alzheimer's dis	ease (moderate)		
Before providing patient identifiable	e data on this form, plea	se confirm that the patient (or ir	the case of a minor or	vulnerable adult v	vith the
parent/legal guardian/carer) has give				to be passed to th	ne CCG and/or
CSU for processing this funding re-	quest and validating sub	sequent invoices. Consent give	en: 🔲		
If there is more than one NICE-app	roved treatment availab	le, a discussion between the re-	sponsible clinician and	the patient has tal	ken place
about the advantages and disadvar					
patient is likely to adhere to treatmand price per dose) unless an orde			osen (taking into accou	nt administration of	costs, dosage
		III the TAS.	Duration		
Patient NHS No:	Trust:		Practice Name:		
Patient	Consultant				
Hospital	Making		Practice Postcode:		
No:	Request:		i ostcode.		
Patient's			Practice		
Initials and DoB:			Code:		
Notification			Contact		
Email	(@	NHS.net account ONLY)	name &		
Address:			number:		
Start date	Provider:	☐Private ☐NHS			
of	Summilian.		Sub-Type:		
requested treatment:		Homecare Hospital		/A 🔽	
			(if applicable)		
By completing this form, you confir commissioning statement. Any req					al
commissioning statement. 7thy req	dests willon fall outside	or this doc will require air maivi	dual randing request (i	110).	
For support regarding IFRs, please	e contact: norfolkicd@nl	ns.net			
For support regarding the criteria li	sted below please con	act: norfolknontariff@nhs.net			
Please indicate whether patie	ent meets the following	ig NICE criteria:		Please tick	
1. The three acetylcholinesteras					
monotherapies are recommended the conditions specified in 1.4 are				☐ Yes ☐ No	
·					
Memantine monotherapy is re	ecommended as an opti	on for managing Alzheimer's dis	sease for people with:		
moderate Alzheimer's disease	who are intolerant of or	have a contraindication to ACh	E inhibitors or	☐Yes ☐No	
severe Alzheimer's disease. Tr	reatment should be und	er the conditions specified in re	commendation 1.5.5		
in the NICE guideline on demen		or the conditions specified in re	oonmendation 1.0.0		
3. This recommendation has been	en undated and renlace	d by recommendation 1.5.5 in the	he NICE quideline on		
dementia.	en upuateu anu repiace	d by recommendation 1.5.5 in the	THE TRICE GUIDEIITIE OIT	☐Yes ☐No	
4. If proporibing on AChE inhibit	or (dononozil, golontom	ing or rivestigming) treatment of	hould normally be		
4. If prescribing an AChE inhibite started with the drug with the love					
per dose once shared care has	started). However, an a	Iternative AChE inhibitor could be	pe prescribed if it is	☐Yes ☐No	
considered appropriate when tal- medical comorbidity, possibility			out adherence,		
3.1					
<ol><li>When using assessment scal professionals should take into a</li></ol>					
difficulties that could affect the r				☐Yes ☐No	
professionals should also be mir	ndful of the need to sec	ure equality of access to treatm			
different ethnic groups, in particu	ular those from different	cultural backgrounds.			
6. When assessing the severity					
should not rely solely on cognition These include:	on scores in circumstar	ices in which it would be inappr	opriate to do so.		
mese moude.					
• if the cognition score is not, or is not by itself, a clinically appropriate tool for assessing the severity of that					

patient's dementia because of the patient's learning difficulties or other disabilities (for example, sensory impairments), linguistic or other communication difficulties or level of education or					
if it is not possible to apply the tool in a language in which the patient is sufficiently fluent for it to be appropriate for assessing the severity of dementia or					
• if there are other similar reasons why using a cognition score, or the score alone, would be inappropriate for assessing the severity of dementia. In such cases healthcare professionals should determine the need for initiation or continuation of treatment by using another appropriate method of assessment.					
7. 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.					
8. The product is being used as described by local commissioning position.					
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:					