FOLLOW-UP VISIT PACKET UNIFORM DATA SET (UDS) VERSION 4.0



Examiner's

Form A4a: ADRD–Specific Treatments

ADRC:		PTID:		Form date:/	/ `	Visit #:	initials:				
Language 1 Englis 2 Spani	sh □1 In-perso ish □2 Remote		Key (remote r	eason): 1=Too cognitively ii 2=Too physically in 3=Homebound or i 4=Refused in-perso 5=Other	npaired nursing home						
dementi clinical of those the of these treatmen lifetime cell. Info develop	as (ADRD) bioma care or an investig at are FDA-appro treatments as pai nt should be include should be include rmation on the ty	rkers, whether a dise national treatment re ved for disease-mod rt of their clinical car aded on both this for red. If available, the Cl pe of treatment can This form should be	ase-modifying trece recived as part of a ification of ADRD s e at the time of clin m and the A4 Medi inicalTrials.gov ide be found via Clinic completed based o	expected to significantly atment that is FDA-apportinical trial. For treatment hould be included on the ical assessment (e.g., the cation form. Participation that is suming a participant interview on participant interview on PAA. Check only on	roved for ADR sents received his form. If the hey are receivi hin in any ADF d into the "spe marized in "Al v and/or co-pe	D and rece as part of participar ing lecaner RD drug tri ecific treat Izheimer's articipant	eived as part of clinical care, only not is receiving one mab infusions), the all over an individual's ment and/or trial" disease drug report. For additional				
1. Has the participant ever been prescribed a treatment or been enrolled in a clinical trial of a treatment expected to modify ADRD biomarkers? 0 No (END FORM HE 1 Yes 9 Unknown (END FORM HE 1 Yes 9 Unknown (END FORM HE 1 Yes 1 Yes											
1a. Since the last UDS visit, is new information available concerning any of the participant's prescribed treatments or clinical trial(s) of a treatment expected to modify ADRD biomarkers?						1 Yes	(END FORM HERE)				
2.	2. Please provide information about the clinical treatment(s) and/or trial(s). INCLUDE ALL HISTORICAL CLINICAL TREATMENT(S) AND/OR TRIAL(S) even if there is no new information on those treatments/trials. If there is a new treatment and/or trial to report, please add it after all of the historical information. (If participant is exposed to more than two treatments and/or trials, use extended table on Page 2):										
Specific Primary Drug Target treatment and/ (check all that apply) or trial		Start date (99/9999 =Unknown)	End date (month/year) (99/9999=Unknown; 88/8888=Ongoing)	How was the treatment provided?		If clinical trial, in which group was the participant?					
1 Amyloid beta 1 Tau 1 Inflammation 1 Synaptic plasticity/ neuroprotection 1 Other target(s)					☐ 1 Clinical care ☐ 2 Clinical trial ☐ 3 Clinical care and clinical trial		1 Active treatment 2 Placebo 9 Unknown				
3. Has the participant ever experienced amyloid related imaging abnormalities-edema (ARIA-E), amyloid related imaging abnormalities-hemorrhage (ARIA-H), or other major adverse events associated with treatments expected to modify ADRD biomarkers?											
3a	experience of related imag	of amyloid related in ging abnormalities–ł	naging abnormalit nemorrhage (ARIA-	ation available concerning the participant's g abnormalities–edema (ARIA-E), amyloid rrhage (ARIA-H), or other major adverse events to modify ADRD biomarkers?			o No (END FORM HERE) 1 Yes 9 Unknown (END FORM HERE)				
3	3b. What major adverse events associated with treatments expecte to modify ADRD biomarkers did the experience? (check all that apply)						3b3. □ 1 Other issues				
	to modify Al	DRD biomarkers did	they 3b2. 1 A	Amyloid related imagin	g	-					

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1 Cummings et al., "Alzheimer's disease drug development pipeline: 2024," Alzheimer's and Dementia. 2024 April 24; 10(2):e12465.

2.	Please provide information about the clinical treatment(s) and/or trial(s). INCLUDE ALL HISTORICAL CLINICAL TREATMENT(S) AND/OR TRIAL(S) even if there is no new information on those treatments/trials. If there is a new treatment and/or trial to report, please add it after all of the historical information. (continued from Page 1):									
	ry Drug Target k all that apply)	Specific treatment and/ or trial	Start date (month/year) (99/9999 =Unknown)	End date (month/year) (99/9999=Unknown; 88/8888=Ongoing)	How was the treatment provided?	If clinical trial, in which group was the participant?				
1 Tau 1 Infl 1 Syr ne	nyloid beta I lammation naptic plasticity/ uroprotection her target(s)	NCT			1 Clinical care 2 Clinical trial 3 Clinical care and clinical trial	1 Active treatment 2 Placebo 9 Unknown				
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_____ Form date: ____ / ____ / ____ Visit #: _

Participant ID: