INITIAL VISIT PACKET UNIFORM DATA SET (UDS) VERSION 4.0



Form D1a: Clinical Syndrome

| ADRC: | PTID: | F | orm date:/ | / | Visit #: | initials: | | | |
|--|--|---------------------|--|---------------------------------------|-----------------------|---|--|--|--|
| Language: 1 Englis 2 Spanis | h □ 1 In-person | Key (remote reas | 2=Too physic | cally impaired and or nursing home | | | | | |
| INSTRUCTIONS: This form is to be completed by the clinician. For additional clarification and examples, see the UDS Coding Guidebook for Form D1a. Check only one box per question. | | | | | | | | | |
| _ | iagnosis method—responses in this form \Box 2 Formal conse | | • | or more clinicians or | other informal gr | oup) | | | |
| Sectio | n 1 – Level of impairment – | Unimpaired co | gnition/beho | avior, SCD, MC | I/MBI, or der | mentia | | | |
| 1 A 2 | 2. Does the participant have: Unimpaired cognition (e.g., cognitive performance and functional status (i.e., CDR) judged to be unimpaired)? AND Unimpaired behavior (i.e., the participant does not exhibit behavior sufficient to diagnose MBI – see MBI section starting at Q7) or dementia due to FTLD or LBD and/or FTLD behavior and language domains=0? No (SKIP TO QUESTION 3) 1 Yes (CONTINUE TO QUESTION 2a) Note: For those with longstanding cognitive impairment that does not represent a decline from their usual functioning, consider checking Question 5b for a diagnosis of "Cognitively Impaired, Not MCI/dementia". | | | | | | | | |
| Subjec | tive Cognitive Decline | | | | | | | | |
| 2a. | Does the participant report 1) signific 2) no neuropsychological evidence of | | | - | 0 No (END FO | ORM HERE) | | | |
| 2b. | As a clinician, are you confident that is clinically meaningful? | :he subjective cogi | nitive decline | 0 No (END FO | | | | | |
| Demer | itia criteria | | | | | | | | |
| Participa | ement #1: ant has cognitive or behavioral (neuro ns that meet <u>all of the following criter</u> | | Requirement Participant m following do | nust have impair | ment in <u>one* c</u> | or more of the | | | |
| Interfere with ability to function as before at work or at usual activities Represent a decline from previous levels of functioning Are not explained by delirium or major psychiatric disorder Include cognitive impairment detected and diagnosed through a combination of: 1) history-taking; 2) objective assessment (bedside or neuropsychological testing) Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired ability to acquire and remember new information of: Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handling of complex tasks, poor judgment Impaired reasoning and handlin | | | | | | nplex tasks, poor nportment guage in PPA, behavior | | | |
| _ | oes the participant meet criteria for denderson to the local of the lo | mentia? | TO QUESTION 6 | 5a) | | | | | |

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| Participan | ID: | Form date: | _ / / | Visit #: | | | | |
|---|--|----------------------------------|------------------------|---|-----------------------------|--|--|--|
| | | | | | | | | |
| Section | n 1 – Level of impairmen | t | | | continued | | | |
| MCI cor | e clinical criteria | | | | | | | |
| Check all | criteria that apply in Q4. | | | | | | | |
| | | | | | | | | |
| If all three | criteria are checked, choose 1=MC | l for Q4b. If less than 3 | criteria are met, cho | ose 0=No for Q4b. | | | | |
| 4b. | Does the participant meet all thre (amnestic or non-amnestic)? | ee of the above crite | ria for MCI | O No (CONTINUE TO QUESTION) 1 Yes (SKIP TO QUESTION) | | | | |
| Cogniti | vely impaired, not MCI/den | nentia | | | | | | |
| | ose of the "Cognitively impaired, n nt or decline who do not meet for | | egory is to capture t | those individuals with evider | nce of cognitive | | | |
| Check all | Check all applicable criteria for cognitively impaired, not MCI/dementia in Q5, using any relevant data. Any conditions contributing to impairment (e.g., substance abuse or medications) should be identified in Section 3. (Note: If recent onset (not longstanding impairment), indicate the cognitive symptom(s) in Form B9 – Clinician Judgment of Symptoms.) | | | | | | | |
| | 5. I Evidence of functional impairment (e.g., CDR SB>0 and/or FAS>0), but available cognitive testing is judged to be normal Cognitive testing is abnormal but no clinical concern or functional decline (e.g., CDR SB=0 and FAS=0) 1 Longstanding cognitive difficulties, not representing a decline from their usual function (e.g., early developmental differences remote TBI, other medical condition with clear effects on cognition) 1 Other (SPECIFY): | | | | | | | |
| | ne criteria in Q5 are met, or if only ia is met in Q4, select 0=No for Q5 | | eria from Q4 are met | , choose 1=Yes for Q5b. Note | e, if <u>only</u> the third | | | |
| 5b. | Does the participant meet any credementia? | iteria for cognitively | impaired, not MCI/ | 0 No (SKIP TO QUESTION 1 Yes (SKIP TO QUESTION | | | | |
| Affecte | d Domains – Dementia and | MCI | | | | | | |
| Choose d | omains that are impaired at the cu abase. | ırrent visit. <u>Select on</u> e | e or more as Impaire | ed; all others will default to u | inimpaired in the | | | |
| Note on behavior changes : For patients with <i>dementia</i> who have behavior changes, record the presence of behavioral changes here (not in the following MBI section) by marking Q6f as Impaired and skipping the MBI section (SKIP TO Q8a). For behavioral changes in the context of an MCI (or as an isolated) symptom, consider a diagnosis of MBI in the next section. | | | | | | | | |
| | | | | | Impaired | | | |
| 6a. | Memory | | | | □ ₁ | | | |
| 6b. | Language | | | | □ ₁ | | | |
| 6c. | Attention | | | | □ ₁ | | | |
| 6d. | Executive | | | | □1 | | | |
| 6e. | Visuospatial | | | | □ ₁ | | | |
| 6f. | Behavioral (for participants with a | lementia only; see MB | l for MCI participants |) | □1 | | | |
| 6g. | Apraxia | | | | □ 1 | | | |

| Section | n 1 – Level of impairment | c | ontin | ued |
|---------------------------------------|--|--|--------------|------------|
| Mild Be | havioral Impairment (MBI) core clinical criteria | | | |
| pers Sym Late Not long Sym Larg mini | cipant, co-participant, or clinician identifies a change in the participant's affect, motivation, thou onality that is clearly different from their usual affect, motivation, thought content, behavior, or possed to be present at least intermittently for the last six months or longer onset (i.e., age > ~50, unless early onset neurodegenerative syndrome is suspected) explained by delirium, other psychiatric disorder by DSM criteria (including recent onset, longsta standing disorder). ptoms interfere with at least one of these: work, interpersonal relationships, social activities ely preserved independence in other functional abilities (no change from prior manner/level of f mal aids or assistance) possible participant meet criteria for MBI? (If participant meets criteria for | oersonality anding or recurr functioning, or | rence d | |
| | ementia an MBI diagnosis is excluded.) | TO QUESTION | 7a) | |
| (N | BI affected domains — <u>Select one or more</u> affected domains ote: If "Yes" is indicated in any domain below, the participant should have a corresponding symptom checked on F Symptoms, either from among the specific symptoms denoted there, or in "other") | orm B9 — Clinicio | an Judg | ment |
| | | | lo | Yes |
| 7a. | Motivation (e.g., apathy symptoms on Form B9) | | <u></u> 0 | 1 |
| 7b. | Affective regulation (e.g., anxiety, irritability, depression, and/or euphoria symptoms on Form B9) | | 0 | <u> </u> |
| 7c. | Impulse control (e.g., obsessions/compulsions, personality change, and/or substance abuse symptoms on For | | 0 | |
| 7d. | Social appropriateness (e.g., disinhibition, personality change, and/or loss of empathy symptoms on Form B | 9) | 0 | |
| 7e. | Thought content/perception (e.g., delusions and/or hallucinations on Form B9) | | 0 | |
| Sectio | n 2 – Clinical syndrome | | | |
| using all a neuropsy section in | pose of Section 2 is to assign a clinical syndrome to participants with dementia and, when approximate clinical, exam, and neuropsychiatric data. This should be done using clinical information in the proximate chological testing, ideally without reference to biomarker data (which is incorporated into the proximal Form D1b). This is not always possible and thus Q9 allows centers to record when biomarker dated the clinical diagnosis. | n and cognitive Etiological Dia | e/ ignose | |
| | plicable syndrome(s) as present; all others will default to Absent in the NACC database. Note that clinical criteria (for instance, this is common for MCI and "impaired, not MCI"). In this case, leave 8 | | nay no | it |
| | | | Pre | sent |
| 8a. | Amnestic predominant syndrome | | |]1 |
| 8b. | Dysexecutive predominant syndrome | | |]1 |
| 8c. | Primary visual presentation (such as posterior cortical atrophy (PCA) syndrome) | | |]1 |
| 8d. | Primary progressive aphasia (PPA) syndrome: | | |] 1 |
| 86 | If present, select one: 1 Logopenic PPA 2 Semantic PPA 3 Nonfluent/agrammatic PPA 4 Primary progressive apraxia of speech 5 PPA other/not otherwise specified | | | |
| 8e. | Behavioral variant frontotemporal (bvFTD) syndrome | | |]1 |
| 8f. | Lewy body syndrome | | |] 1 |
| 8 | f1. If present, select one: 1 Dementia with Lewy bodies 2 Parkinson's disease 3 Parkinson's disease dementia syndrome | | | |
| 8g. | Non-amnestic multidomain syndrome, not PCA, PPA, bvFTD, or DLB syndrome | | |]1 |

Form date: ____ / ____ / ____ Visit #: __

Participant ID:

| Partici | pant ID: | : Form date: | / | _ / | | Visit #: | |
|--|--|---|--|--|---|---|---|
| Section 2 – Clinical syndrome continued | | | | | | | |
| | | | | | | | Present |
| 8 | 3h. Pr | rimary supranuclear palsy (PSP) syndrome | | | | | □ ₁ |
| | 8h1. If present, select one: 1 Richardson's syndrome criteria 2 Non-Richardson's | | | | | | |
| | 8i. Tr | Traumatic encephalopathy syndrome | | | | | |
| | 8j. Co | Corticobasal syndrome (CBS) | | | | | |
| 8 | Bk. M | lultiple system atrophy (MSA) syndrome | | | | | 1 |
| | 8k1. | If present, select one: 1 MSA-predominant cerebellar ataxia (MSA-C) 2 MSA-predominant Parkinsonism (MSA-P) 3 MSA-predominant dysautonomia | | | | | |
| | 8I. O | ther (SPECIFY): | <u> </u> | | | | □ ₁ |
| 9. | | ate the source(s) of information used to assign the cli t one or more as Yes ; all others will default to No in th | - | | | | |
| | | | | | | | Yes |
| ġ | 9a. Cl | linical information (history, CDR) | | | | | □ ₁ |
| 9 | 9b. Cognitive testing | | | | | | □ ₁ |
| 9 | 9c. Bi | iomarkers (MRI, PET, CSF, plasma) | | | | | □ ₁ |
| Section 3 – Primary or contributing non-neurodegenerative or non-CVD conditions | | | | | | | |
| Sect | tion 3 | 3 – Primary or contributing non-neuro | degene | erative | or non- | CVD conditi | ions |
| The p | urpose nust be | B – Primary or contributing non-neuro of Section 3 is to identify conditions or disorders that filled out for those with cognitive or behavioral impa a primary, contributing, or non-contributing cause of | t are prese airment (i.e | nt and po ., MCI, M | otentially co BI, dementia | ntributing to the a, etc.) Indicate v | e clinical syndrome. whether a given |
| The properties of the properti | urpose nust be tion is a | of Section 3 is to identify conditions or disorders that filled out for those with cognitive or behavioral imparations a primary, contributing, or non-contributing cause of r more syndrome(s) as Present ; all others will default | t are present firment (i.e the observ | nt and po ., MCI, M ved impa | otentially co BI, dementia irment, basa | ntributing to the a, etc.) Indicate v ed on the clinicia | e clinical syndrome. whether a given an's best judgment. |
| The portion of the po | urpose nust be tion is a t one of ed as 1 der to d | of Section 3 is to identify conditions or disorders that filled out for those with cognitive or behavioral impa a primary, contributing, or non-contributing cause of | t are present hirment (i.e the observe to Absent in | nt and po ., MCI, M ved impa in the NA cause cli | otentially co BI, dementia irment, base ICC databas nically signi | ntributing to the a, etc.) Indicate v ed on the clinicia e. Only one diago ficant distress or | e clinical syndrome. whether a given an's best judgment. mosis should be impairment in |
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| The portion of the po | urpose nust be tion is a t one or ed as 1 der to o , occup | of Section 3 is to identify conditions or disorders that filled out for those with cognitive or behavioral imparts a primary, contributing, or non-contributing cause of r more syndrome(s) as Present ; all others will default = Primary . diagnose a disorder, DSM-5-TR criteria require that stational, or other important areas of functioning. For the | t are presen airment (i.e the observ to Absent i symptoms more guida | nt and po ., MCI, M ved impa in the NA cause cli | otentially co BI, dementia irment, base CC databas nically signi the UDS Co | ntributing to the a, etc.) Indicate v ed on the clinicia e. <i>Only one diagi</i> ficant distress or ding Guidebool | e clinical syndrome. whether a given an's best judgment. nosis should be impairment in c, Form D1a. |
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| The p This m condii Selecte *In ore social, 10. 11. 12. | Major Other Anxie If | e of Section 3 is to identify conditions or disorders that a filled out for those with cognitive or behavioral impartant primary, contributing, or non-contributing cause of a more syndrome(s) as Present; all others will default = Primary. diagnose a disorder, DSM-5-TR criteria require that solutional, or other important areas of functioning. For a condition redepressive disorder (DSM-5-TR criteria*) respecified depressive disorder (DSM-5-TR criteria*) are disorder (DSM-5-TR criteria*) ophrenia or other psychotic disorder (DSM-5-TR ia*) ety disorder (DSM-5-TR criteria*) present, (SPECIFY) (check all that apply): 1 Generalized anxiety disorder 1 Panic disorder 1 Obsessive-compulsive disorder (OCD) | t are present in the observation of the observation | nt and po ., MCI, M ved impa in the NA cause cli ance see 10a. 11a. 12a. | otentially co BI, demential irment, base CC databas nically signithe UDS Co Primary 1 1 1 1 | ntributing to the a, etc.) Indicate ved on the cliniciae. Only one diagrams or ding Guidebook Contributing 2 2 2 2 | e clinical syndrome. whether a given an's best judgment. mosis should be impairment in c, Form D1a. Non-contributing 3 3 3 3 |

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| Section 3 – Primary or contributing non-degenerative or non-CVD conditions continued | | | | | | | |
|--|-----------------|--|----------------|------|----------------|--------------|-----------------------|
| | | Condition | Present | | Primary | Contributing | Non-contributing |
| 16. | specti | opmental neuropsychiatric disorders (e.g., autism rum disorder (ASD), attention-deficit hyperactivity ler (ADHD), dyslexia) | □ 1 | 16a. | □ 1 | \square_2 | <u></u> 3 |
| 17. | Deliriu | um (DSM-5-TR criteria*) | □ 1 | 17a. | 1 | _2 | 3 |
| 18. | Other | psychiatric disorder (DSM-5-TR criteria*) | □ 1 | 18a. | □ 1 | \square_2 | 3 |
| | 18b. | If present, (SPECIFY) : | | | | | |
| 19. | (Distin | natic brain injury nct from TES and CTE, which are documented as a al Syndrome and Etiologic Diagnosis, respectively) | □ 1 | 19a. | □ 1 | \square_2 | □ ₃ |
| 20. | Epilep | osy | 1 | 20a. | □ 1 | 2 | 3 |
| 21. | Norm | al-pressure hydrocephalus | □ 1 | 21a. | 1 | _2 | 3 |
| 22. | CNS N | leoplasm | □ 1 | 22a. | □ 1 | \square_2 | 3 |
| 22b. If present, select one: 1 Benign 2 Malignant | | | | | | | |
| 23. | Huma | n immunodeficiency virus (HIV) infection | 1 | 23a. | □ 1 | \square_2 | 3 |
| 24. | Post C | OVID-19 cognitive impairment | □ 1 | 24a. | □ 1 | \square_2 | 3 |
| 25. | Sleep apnea) | apnea (i.e., obstructive, central, mixed or complex sleep | | 25a. | □ 1 | <u></u> | □3 |
| 26. | infect | tive impairment due to other neurologic, genetic, ious conditions (<i>not listed above</i>), or systemic se/medical illness (as indicated on Form A5/D2) | □ 1 | 26a. | □1 | <u>2</u> | 3 |
| 26 | ib. If p | oresent, (SPECIFY): | | | | | |
| 27. | Cogni | tive impairment due to alcohol use or abuse | □ ₁ | 27a. | □ ₁ | _2 | 3 |
| 28. | Cogni | tive impairment due to substance use or abuse | □ 1 | 28a. | □ ₁ | \square_2 | 3 |
| 29. | Cogni | tive impairment due to medications | □ ₁ | 29a. | □ ₁ | \square_2 | 3 |
| 30. | Cogni | tive impairment not otherwise specified (NOS) | □ ₁ | 30a. | □ ₁ | \square_2 | 3 |
| 30b. If present, (SPECIFY): | | | | | | | |
| 31. | Cogni | tive impairment not otherwise specified (NOS) | 1 | 31a. | □ 1 | \square_2 | 3 |
| 31 | b. If | present, (SPECIFY): | | | | | |
| 32. | Cogni | tive impairment not otherwise specified (NOS) | 1 | 32a. | □ 1 | \square_2 | 3 |
| 32 | h If | present (SDECIEV). | | | | | |