The Utility of the Mild Behavioral Impairment-Checklist in Detecting Neuropsychiatric Symptoms in Mild Cognitive Impairment and Dementia

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Introduction:
Dementia is one of the most common neurological disorders globally with cases expected to double by 2031 in Canada alone. Although memory loss is a hallmark symptom, neuropsychiatric symptoms such as anxiety and agitation are early markers. Mild Behavioral Impairment (MBI) is an at-risk state for dementia characterized by sustained neuropsychiatric symptoms. It is a develop that portrays the rise at ≥50 years old of supported and effective neuropsychiatric indications (NPS), as a forerunner to intellectual decrease and dementia. MBI portrays NPS of any seriousness, which are not caught by customary mental nosology, persevere for in any event a half year and happen ahead of time of or working together with gentle psychological impedance. While the depiction and picture of MBI has been operationalized in the investigates demonstrative measures, there is no instrument that precisely reflects MBI as portrayed. We built up the MBI Checklist (MBI-C) to survey inspiration, disposition, motivation control, social fittingness and discernment in pre-dementia patients.

Methods:
The MBI-C has been administered in the Cognitive Neuroscience Clinic at the University of Calgary, Canada (n=227). We analyzed baseline MBI-C and gold standard neuropsychiatric inventory questionnaire (NPI-Q) scores in relation to MoCA scores in normal cognition (n=38), mild cognitive impairment (n=93) and dementia (n=74) patients using linear regression. An iterative procedure guaranteed things mirrored the five MBI spaces of 1) diminished inspiration 2) enthusiastic dysregulation 3) motivation dyscontrol 4) social impropriety and 5) irregular observation or thought content. Instrument language was built up from the earlier to relate to non-maniacal practically free more seasoned grown-ups.

Results:
With increasing severity of cognitive diagnosis, neuropsychiatric symptoms worsen (MBI-C and NPI-Q scores increase) and cognition declines (MoCA score decreases). Those with worsened cognition tend to be older, female and have less education. We found for every one point increase in MBI-C score, there is a 0.082 point decrease in MoCA score (p=0.007). For every one point increase in NPI-Q score, there is a 0.192 point decrease in MoCA score (p=0.006). There is no alteration yet age and training are confounders.

Discussion:
In the psychiatric outpatient clinic, the popularity of MBI was 3.5% and the frequency rate of dementia was 30.7 cases per 1000 person-years. MBI, MCI without MBI, and SCD without MBI increased the risk of dementia, while sleep disorder and depressive episode did not. In the MCI patients, those with affective dysregulation tended to develop dementia with a hazard ratio of 1.646 compared to those without. Therefore, MBI, especially affective dysregulation domain, might be associated with dementia.

Conclusion:
Given that the MBI-C is more sensitive in detecting neuropsychiatric symptoms in pre-dementia populations, there is a shallower point change in MoCA score. The MBI-C may be used to detect neuropsychiatric symptoms in normal cognition and MCI patients. Both cognitive and behavioral scales should be used to assess neuropsychiatric symptoms and cognitive decline in patients. Studies are required to determine the prognostic value of MBI for dementia development and for predicting different dementia subtypes.