The early detection of cognitive impairment: Two approaches to assess longitudinal trajectories

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Today’s Presentation

Briefly introduce two methods suited for the analysis of longitudinal trajectories.

1) Change point model
2) Latent (Cluster) Trajectory model
Why Trajectories?

Early detection of dementia is becoming more important for treatment and long-term planning.

Therefore, it is critical to detect early signals indicating the transition from normal cognitive aging to MCI and early dementia.

Studying the trajectories of biomarkers and cognitive and functional measurements could help identify these “early signals”.


This study found that approximately 5 years before the diagnosis of dementia, memory function (as measured by Buschke Selective Reminding test) started accelerating in decline. (Sample: Bronx Aging Study)
When does acceleration (change point in slope) starts?

\[
\text{Outcome}_{it} (\text{memory test score}) = \text{intercept} + (\text{age}_{it} - 75) + (\text{age}_{it} - (\text{aged}_i - \tau))
\]

\[
(\text{age}_{it} - (\text{aged}_i - \tau)) = \max (0, \text{age}_{it} - (\text{aged}_i - \tau))
\]
Research questions

What variables are sensitive to transition from normal cognition to MCI?

For example…..

*When* do ventricular volumes change; Or neuropsychological test scores; Or motor function (gait speed) change?
Application of Change Point Analysis (Examples)

1) Ventricular volume
2) Neuropsychological tests
3) Gait speed
4) Hand tapping speed

DATA: The Oregon Brain Aging Study (aka OBAS) - a longitudinal community-based cohort of healthy elderly.

At entry, all subjects were cognitively intact with CDR=0 and a MMSE score $\geq 24$. 
Application of Change Point Analysis

Step 1: Mixed effects model
Assess if overall trajectories were different between normal cognition vs. those who developed MCI

Step 2: Among those who developed MCI, determine whether there was a change point, and if so, when the change point occurred in relation to the development of MCI.
When the acceleration starts in relation to the development of MCI?

Ventricular volume

When the acceleration starts?
Neuropsychological tests

Logical Memory I

Logical Memory II

Category Fluency

Block Design

When the acceleration starts in relation to MCI?

Gait Speed

14.2 years

6.0 yrs

Mean age at MCI conversion 89.9

Buracchio, et al., presented at the 2009 AAN Annual Meeting in Seattle
When the acceleration starts in relation to the development of MCI?

**Tapping Speed (Non-Dominant Hand)**

![Graph showing tapping speed over age](image)

- **Mean age at MCI conversion:** 89.9 years
- **Tapping speed for men:** 6.16 years
- **Tapping speed for women:** 0.41 years

The graph illustrates the decline in tapping speed as a function of age, distinguishing between men and women. The vertical line indicates the mean age at MCI conversion, which is consistent with the observed tapping speed decrease.
Conclusion

• Change points were recognized several years prior to the development of MCI in several domains (brain volume, cognition, motor function).

• These change points may be useful markers for guiding the timing of assessments used in clinical trials and other prodromal dementia research designs.
Limitations of Change Point Analysis

• The amount of acceleration is “on average”. It is hard to generalize the result to individuals in clinical practice (e.g., decline in walking speed by 0.023 meter/second/year--predicts MCI a decade later?).

• Ecological fallacy: Using group data, we found the order of accelerations (gait speed->ventricular volume->neuropsychological tests->hand tapping speed). But individuals might not necessarily follow this order.

Further steps are required to translate the findings into clinical practice for early detection of MCI/Dementia
2) Latent Trajectory Analysis

• **Latent Trajectory Analysis**

• Implemented in **PROC TRAJ** procedure in SAS (Jones BL, et al., 2001)
  [http://www.andrew.cmu.edu/user/bjones](http://www.andrew.cmu.edu/user/bjones)
A latent class analysis which identifies homogeneous trajectory patterns and associated factors for each pattern. (Nagin, 1999).

Given that there are K latent trajectory groups, the conditional distribution of the observable outcome for subject i (y_i), given risk factors z_i, is written as follows:

\[
\Pr(\mathbf{y}_i | \mathbf{z}_i) = \sum_{k=1}^{K} \Pr(C_i = k | Z_i = z_i) \Pr(Y_i = y_i | C_i = k)
\]

where C_i is latent group identification for subject i.
Application of Latent Trajectory Analysis to an Epidemiological Study

Application of Latent Trajectory Analysis

ORCATECH (Oregon Center for Aging and Technology):

BRP Study - Intelligent Systems to Assess Aging Change (aka ISAAC study).

Following approximately 230 Portland area elders (mean age, 84) with in-home monitoring technologies
Motion Sensors
Contact Sensors
Ekahau ID tag
Community-wide home-based assessment: “The ORCATECH Living Laboratory”
Intelligent Systems to Assess Aging Change (aka ISAAC study)

Goal: to determine whether the unobtrusive monitoring data from activity and computer sensors can be used to effectively predict when an elder starts to develop cognitive decline.
Application of Trajectory Analysis to walking speed measured unobtrusively at home

Data Source: Intelligent Systems to Assess Aging Change (ISAAC)
The Oregon Center for Aging and Technology (ORCATECH)
Summary

• **Analytical methods** aimed to **efficiently examine trajectories** of various predictive variables for the development of MCI
  - Change point model (Hall CB, et al., 2000)
  - Latent Trajectory model (Jones BL, et al., 2001)

• **Allow us to clarify what changes are occurring during a (long) pre-symptomatic period of dementia**

• **Further methodological developments are required to translate these findings into clinical practice for the early detection of MCI/dementia onset.**
Thank you!