Novel Regulatory Science Research on Drugs for Alzheimer's Disease

-A Guideline on the Clinical Evaluation of Drugs for Alzheimer’s Disease (Interim Report)-

Takashi Moritoyo M.D., Ph.D.

Pharmaceuticals and Medical Devices Agency

The University of Tokyo Hospital
Disclosure of conflict of interest

There are no companies, etc. in a relation of conflict of interest requiring disclosure in relation to the presentation.

Disclaimer

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MHLW launched a project termed “Accelerating regulatory science initiatives” to promote the development of innovative drugs and its approval review in 2012.
Academic Institutions Involving in the Project

Pharmaceutical Area
Medical Device Area
Regenerative Medicine Area

Newly joined in 2013

Alzheimer’s disease

24 institutes in total (as of May, 2013)
21 research institutes participated in this program in 2012 and 18 researchers were accepted as specially appointed experts (incl. part-time) while 30 (*) reviewers were dispatched (incl. part-time). Three more institutes joined in 2013. (*) excluding those dispatched as instructors

U. Tokyo Hospital


U. Tokyo Grad. Sch. of Eng.
Regulatory science research on the drug development for Alzheimer's disease

【Objective】
Establish standards for clinical evaluation of drugs for Alzheimer's disease (AD)

【Methods】
- Simulations and analyses on histories obtained from ADNI and clinical trials

【Goals】
- Facilitate development of new anti-AD drugs
- Facilitate establishment of standards for evaluation of new anti-AD drugs
(1) Establishment of Standards for Clinical Evaluation of Anti-AD Drugs using Biomarkers

AD Neuroimaging Initiative

Neuroimaging (PET・MRI)
Biomarker (peptide levels)
Cognitive function test

Clinical trial, simulations, and etc.

Disease-Modifying Treatment

Biomarker Effects

Biomarker Outcomes

Same Mechanism of Action

Correlation

Clinical Effects

Clinical Trial Outcomes

Establish standards (guideline) for evaluation of new anti-AD drugs

Support development of innovative drugs

Dubois (2010)    AD pathology    Prodromal AD    AD dementia

NIA/AA (2011)    Preclinical AD    MCI due to AD    AD dementia

(Sperling RA, et al., Alzheimers Demnt, 2011)

(Cummings JL, Neurobiol Aging, 2012)
(2) Extension of M & S for prediction of therapeutic benefit of anti-AD drugs

AD Neuroimaging Initiative
Neuroimaging (PET・MRI)
Biomarkers (peptide levels)
Cognitive function test

Developmental process of AD from appearance of amyloid β, then neuronal injury, volumetric change of brain, and finally to dementia

Extension of Modeling & Simulation (M&S)

Disease model
- Biology
  - Natural progression
  - Biomarker(s) / Imaging
  - Genetic effect(s)

Drug model
- Pharmacology
  - Mechanism of action
  - Effectiveness
  - Safety

Trial model
- Pharmacometrics
  - Patient population
  - Method validation
  - Statistic significance

M & S technique developed by this project will be applied to the POC clinical trials of the new anti-AD drug currently being developed by the University of Tokyo.

Robustness of the model will be improved by applying outcomes of various clinical trials including overseas.

(Gobburu JV, et al., Annu Rev Pharmacol, 2009)
The University of Tokyo Hospital

Project Team for Establishment of Standards for Clinical Evaluation of Anti-AD Drugs

Biomerker/ Clinical Evaluation Group
Unit for Early/Exploratory Clinical Development

Clinical Research Support Center

Modeling & Simulation Group
Pharmaceutical Dept.

Establish the clinical evaluation guideline
Support development of new AD drugs

Academic Societies
- Japan Society for Dementia Research
- Societas Neurologica Japonica
- The Japanese Society of Psychiatry and Neurology
- The Japan Geriatrics Society
- Japanese Society of Neurological Therapeutics
- Japanese Psychogeriatric Society

Collaboration

Human Resources Exchange

Using Open Data
- ADNI Clinical trials, etc
- Neuroimaging (PET, MRI)
- Biomarkers
- Cognitive function test

Harmonization

Structure

Office of New Drugs II
Office of Regulatory Science
Omics Project Group
New Statistics Project Group

Japan Society for Dementia Research
Societas Neurologica Japonica
The Japanese Society of Psychiatry and Neurology
The Japan Geriatrics Society
Japanese Society of Neurological Therapeutics
Japanese Psychogeriatric Society
Expected Outcome

- Propose reasonable standards for clinical evaluation of new anti-AD drugs worldwide
- Establish Japanese guideline and facilitate development of anti-AD drugs
“Issues to Consider in the Clinical Evaluation and Development of Drugs for Alzheimer’s Disease” (Interim Report)

“Interim Report” was released to the public in November 2013.

Many comments have been submitted from industry and academia in Japan.
Outline

- Inclusion Criteria
- Efficacy Endpoint
- Clinical Development in Japan
- Preclinical AD
Inclusion Criteria

Efficacy Endpoint

Clinical Development in Japan

Preclinical AD
We recommend to use biomarkers

- To exclude patients who don’t have AD pathology.

(In case of pre-dementia stage) To administer drugs only to patients who have higher risk of developing dementia.
However, further validation and standardization are needed.

- Measurement variability
- Differences between Japanese and non-Japanese are unclear (e.g. cut-off value).

Further investigations in Japanese population are needed.

- J-ADNI
- Biomarker information in clinical trials
Basically, the biomarkers used to select subjects in clinical trials should be available in clinical practice in the same way.

Description in drug labeling about the necessity of measuring biomarkers before administering the drug would be determined based on the risk/benefit of the drug and the medical practices at the time of NDA.
Outline

- Inclusion Criteria
- Efficacy Endpoint
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- Preclinical AD
**Efficacy Endpoint**

- AD dementia
  - Co-primary endpoints are required.
    - Cognition

Activities of daily living

or

Global assessment
Pre-dementia stage of AD (e.g. MCI due to AD)

- Time to a diagnosis of dementia
  - Sufficient training for clinical investigators is necessary to reduce variability.

- Composite scale of cognition and function
  - Clinical meaningfulness must to be explained.
  - Efficacy of the drug should be supported by the result of the secondary endpoint.
    (e.g. separate scales, time to event)
The relationship between biomarkers and clinical effects induced by drug intervention has not been fully understood yet.

At present, it is recommended to evaluate biomarkers as secondary endpoint.

Information of biomarkers is also important to obtain disease modifying claim.
Outline

- Inclusion Criteria
- Efficacy Endpoint
- Clinical Development in Japan
- Preclinical AD
Most of drugs for Alzheimer’s disease have been developed by global clinical trials.

Basically, Phase I study to investigate safety and pharmacokinetics is required in Japanese population.

- Investigation of ethnic differences
- Biomarker information
Phase II study to explore dose-response relationship in Japanese population is required in principle.

- Participate in global phase II study
- Conduct domestic phase II study before global Phase III study

Basically, dose-response relationship should be investigated on the basis of the clinical symptoms not on biomarkers, at present.
Pre-dementia stages of Alzheimer’s disease

- Need to consider the feasibility to conduct clinical trials in Japan.

- If dose-response relationships could be investigated and robust efficacy could be demonstrated within one large scale and long-term clinical trial (e.g. seamless phase II/III study), conducting another phase III study may not be required.
Global phase III study should have a design, in which consistency of the result in primary endpoint can be obtained between Japanese population and entire population.

Outline

- Inclusion Criteria
- Efficacy Endpoint
- Clinical Development in Japan
- Preclinical AD
Further discussion is needed

- Approval System
- Feasibility to conduct long-term placebo control study as post-approval study

J-ADNI2 project, which investigate natural course of preclinical AD, has been just started.
Conclusion

- Using Biomarkers is recommended
  - Patient selection
  - Secondary endpoint
- Further validation is needed
  - Especially in Japanese population
- Increasing global clinical trial
  - Participating in global study from early stage of development is recommended
- We would like to continue the discussion with industry, academia and other regulatory authorities.
Thank you for your attention!