Testing whether the progression of Alzheimer's disease change with the year of publication, additional design and geographical area: a modeling analysis of literature aggregate data

ningyuan zhang  
Shanghai University of traditional Chinese Medicine  
https://orcid.org/0000-0002-9376-2658

Xijun Zheng  
Shanghai University of Traditional Chinese Medicine

Hongxia Liu  
Shanghai University of Traditional Chinese Medicine

Qingshan Zheng (✉ qingshan.zheng@drugchina.net)  
Shanghai University of Traditional Chinese Medicine

Lujin Li  
Shanghai University of Traditional Chinese Medicine

Research

Keywords: Alzheimer's disease, disease progression model, model-based meta-analysis, modelling and simulation

Posted Date: May 7th, 2020

DOI: https://doi.org/10.21203/rs.3.rs-17692/v2

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Version of Record: A version of this preprint was published at Alzheimer's Research and Therapy on May 26th, 2020. See the published version at https://doi.org/10.1186/s13195-020-00630-5.
Abstract

Background Our objective was to develop a disease progression model for cognitive decline in Alzheimer’s disease (AD) and to determine whether disease progression of AD is related to the year of publication, add-on trial design, and geographical regions.

Methods Placebo-controlled randomized AD clinical trials were systemically searched in public databases. Longitudinal placebo response (mean change from baseline in the cognitive subscale of the Alzheimer’s Disease Assessment Scale [ADAS-cog]) and the corresponding demographic information were extracted to establish a disease progression model. Covariate screening and subgroup analyses were performed to identify potential factors affecting the disease progression rate.

Results A total of 134 publications (140 trials) were included in this model-based meta-analysis. The typical disease progression rate was 5.82 points per year. The baseline ADAS-cog score was included in the final model using an inverse-U type function. Age was found to be negatively correlated with disease progression rate. After correcting the baseline ADAS-cog score and the age effect, no significant difference in disease progression rate was found between trials published before and after 2008, and between trials using add-on design and those that did not use add-on design. However, a significant difference was found among different trial regions. Trials in East Asian countries showed the slowest decline rate and the largest placebo effect.

Conclusions Our model successfully quantified AD disease progression by integrating baseline ADAS-cog score and age as important predictors. These factors and geographic location should be considered when optimizing future trial designs and conducting indirect comparisons of clinical outcomes.

Background

Alzheimer’s disease (AD) is a degenerative neurological disease characterized by gradual memory loss and cognitive deterioration. Current FDA-approved AD drugs include cholinesterase inhibitors (CHEIs) and memantine. However, these drugs only provide symptomatic improvement and are unable to reverse cognitive decline in patients with AD [1-4]. Discovery of disease-modifying AD agents is urgent, yet none of the tested drugs showed statistical superior effects compared to placebos in large clinical trials [5-9]. From the perspective of new drug development, abnormal placebo decline might result in an increased risk of false-positive or false-negative outcomes. Therefore, to ensure a correct interpretation of the trial outcomes, it is necessary to predict or evaluate the cognitive decline in placebo-treated AD patients [7, 10].

The disease progression model, which describes the disease trajectory and underlying placebo effect, is widely used in AD clinical research to understand natural AD progression and inform future trial design [11, 12]. To date, several AD disease progression models for various outcomes have been published [11, 13-20]. However, as a large amount of new trial data has been published during the past decade, it is necessary to examine whether the disease progression rate has changed over such a long period. In
addition, we noticed that subjects were usually allowed to continue their treatment of CHEIs or/and memantine as background therapy in recent trials, which is a major distinction from the past trials [21-23]. Another major feature of AD clinical trials in recent years is their multi-nationality. Due to heterogeneities in cultures, access to healthcare, and prevalence of risk factors in different regions, the disease progression rate may differ between countries. However, to our knowledge, only one study has assessed AD progression across geographic regions [24]. They found that AD progression may differ across geographic regions, however, the heterogeneity across patient populations at baseline was not fully considered. Thus, these results need to be confirmed by more extensive data.

In this study, we re-retrieved AD trials and extracted ADAS-cog data from placebo groups to establish a summary-level AD disease progression model. The main objectives of our study were to (1) update the disease progression model, (2) investigate a potential correlation between disease progression and publication year, (3) investigate whether and to what extent disease progression differs between AD trials using or not using standard background therapy, and (4) explore the differences in disease progression in different geographic regions.

**Method**

**Data collection**

The public medical databases PubMed and Embase were searched for AD clinical trials on July 13th, 2019. Key items were classified as "Alzheimer's disease," "clinical trial," "placebo," and "ADAS-cog". Detailed information about the search strategy is provided in the supplementary materials.

Only clinical trials satisfying the following criteria were included in the analysis: (1) enrolled subjects must be diagnosed with dementia caused by AD, (2) trials must use parallel design and blind methods with placebo as the control group, (3) trials must report 11 items on the ADAS-cog scale as the primary or secondary outcome, (4) the sample size of the placebo group should be greater than ten, and (5) the language of the publication must be English.

The following information was extracted from the placebo group of the included AD studies to generate a database:

(1) Characteristics of the publication and trial design, including title, main author, published year, clinical trial registration number, geographic region, trial length, randomized sample size, completed numbers, number of study sites, name of the test drug, whether subjects were allowed administration of standard background therapy (CHEIs or/and memantine) before entering and during the trial, and whether the study used add-on design.

(2) Baseline demographic information such as stage of disease, age, proportion of males, proportion of whites, proportion of patients with background therapy during the trial, proportion of APOE ε4 carriers,
ADAS-cog score, Mini-mental State Examination (MMSE) score, neuropsychiatric inventory (NPI) score, AD duration, and length of education.

(3) Primary outcomes: mean change from baseline in the ADAS-cog score at each reported time point, the corresponding sample size and analyzed dataset (intention to treat, per protocol dataset, etc.), imputation method of missing data (observed case OC or last observation carry forward LOCF). OC data were extracted prior to LOCF data if both were reported. GetData Graph Digitizer (version 2.26.0.20) was used to read longitudinal data if presented graphically.

Disease progression model

Basic model

NONMEN 7.3 (Level 1.0, ICON Development Solutions, USA) was used to fit the time course of AD placebo response data. Model parameters were estimated by the first-order conditional estimation method with interaction (FOCE-I).

Placebo response, which consisted of both disease progression and the theoretical placebo effect, was defined as the mean change in ADAS-cog from baseline in the placebo group [11, 19, 25]. Disease progression refers to the natural change of cognitive function for AD subjects without any intervention, while the component of the placebo effect typically describes the temporary improvement due to the subjects’ expectations after entering the study. The structural model of the placebo response is shown in Equation 1, where $\alpha$ is the slope parameter used to define the rate of linear disease progression and $t$ is the time course of the placebo effect.

[Please see the supplementary files section to view the equation.] (1)

The mathematical expression of the placebo effect models is shown in equations 2, where $\beta$ represents the maximum extent of the placebo effect and $t_{1/2}$ is the time required to reach half of the maximum placebo effect. Besides, sigmoid Emax, inverse Bateman, and an exponential model are also explored to describe the longitudinal placebo effect if possible.

[Please see the supplementary files section to view the equation.] (2)
In AD trials, the disease status of cognitive function might either worsen or improve for different subjects. In addition, the nocebo effect may also affect cognitive function. Hence, inter-study variability for $\alpha$ and $\beta$ was modeled using a proportional error model to allow individual parameter estimates to be either positive or negative (Equation 3). For other structure model parameters, an exponential error model was used because their values were always positive (Equation 4). In equations 3 and 4, $P_i$ is the individual parameter estimate of the $i^{th}$ study and $P_{pop}$ is the population parameter estimate. $\eta_i$ is the random effect for the $i^{th}$ study to characterize the deviation between $P_i$ and $P_{pop}$, which was assumed to follow a normal distribution with a mean of 0 and variance of $\omega^2$. The residual error $\epsilon$ between the measured observation $Y$ and the individual prediction $F$ was modeled using an additive model and then weighted by the inverse of the square root of the corresponding sample size $N$ (Equation 5). $\epsilon$ was assumed to be normally distributed, with a mean of 0 and variance of $\delta^2$.

[Please see the supplementary files section to view the equation.] (3)

[Please see the supplementary files section to view the equation.] (4)

[Please see the supplementary files section to view the equation.] (5)

**Covariate screening**

The correlation between the inter-study variability of basic model parameters and the candidate covariates was explored after establishment of the basic model. The dichotomous covariates, such as whether using add-on design, were tested on the structure model parameters by introducing indication variables 0 and 1. Continuous covariates were tested using a linear equation or power function after normalization of the median value. In particular, the ceiling effect of the ADAS-cog scale should be noted when examining the baseline ADAS-cog effect on the rate of disease progression. When possible, the inverse U-type function was used to describe the nonlinear relationship between ADAS-cog effect and the rate of disease progression [13, 16], as shown in Equation 6, where $\alpha_{typical}$ is the typical population estimate of $\alpha$, $\text{ADAS}_{baseline}$ is the baseline ADAS-cog score, $\text{ADAS}_{median}$ is the median value among the included studies, and $\theta$ is the correction factor to control the shape of the curve.

[Please see the supplementary files section to view the equation.] (6)
Covariate screening was performed in a stepwise manner. Evaluation of the covariate was based on the change value in the objective function value (OFV) using the likelihood ratio test method, decrease in inter-study variability, and prior knowledge of plausibility. In the forward inclusion step, a reduction in OFV of more than 3.84 points ($\chi^2, P < 0.05, df = 1$) was used as the criterion. Upon establishing the full model, the included covariates were removed successively to evaluate their necessity. An increase in OFV of more than 6.64 ($\chi^2, P < 0.01, df = 1$) was considered statistically significant to retain this covariate in the final model.

**Model evaluation**

Three methods were used to evaluate the model quality as follows [26]. (1) The diagnostic graphs were used to assess the consistency between the model prediction and the observation and inspect the randomness of the residual error distribution. (2) A non-parametric bootstrap method was used to generate 1,000 new datasets by random sampling with replacement from the original dataset and using the identical model structure to fit and re-estimate the model parameters. The final model parameter estimates and their 95% confidence intervals (CI) were then compared with the median and 95% quantile of the bootstrap replicates to verify the precision and stability of the final model estimation. (3) Monte Carlo simulations were performed on 1,000 template datasets based on the final model parameter estimates. Through a visual predictive check, the median and the 95% prediction interval of the simulated time course of ADAS-cog were plotted and overlapped with the observed data to qualify the predictive performance of the final model.

**Subgroup analyses**

We conducted subgroup analyses of disease progression rate ($\alpha$) and maximum placebo effect ($\beta$) with regard to geographic regions, trial design (add-on design or not), and publication year (before or after 2008). The analysis method included two main steps. The first step was to obtain the estimated values of model parameters and their standard errors of each study [27]. The original individual estimates of model parameters were then adjusted by inverse calculation of covariate functions, which aimed to deduce the varying degrees of covariate impact across different studies [28, 29]. In the second step, the estimated model parameters were pooled under these predefined subgroups using a random effect meta-analysis of a single mean to obtain the overall mean and the 95% CI.

**Results**

**Characteristics of the included studies**
Finally, 134 articles, which involved 140 independent clinical trials and 19,210 placebo subjects, were eligible for inclusion. The screening flow chart and the reference list are shown in Figure 1 and Additional file. The total number of ADAS-cog observations was 455. The median value (minimum–maximum) of baseline age, ADAS-cog score, and MMSE score were 73.5 years (58.0–81.7), 24.5 points (13.1–39.3), and 19.4 (13.2–25.4) points, respectively. The number of studies using the add-on design was 27, all of which were published after 2008 (Additional file Figure 1). The baseline demographic characteristics were similar as in those studies not using the add-on design. Baseline ADAS-cog scores for trials published after 2008 were significantly lower than those published before 2008 (-2.56, 95 % CI: -3.96, -1.15). The other baseline demographics were comparable between the two groups (Table 1).

The countries where trials were conducted were combined into regions based on ethnicity and geography. Regions were as follows: North America (USA and Canada), Europe/Oceania (UK, France, Germany, Italy, the Netherlands, Norway, Poland, Russia, Ukraine, Spain, Sweden, Australia, and New Zealand), East Asia (China, Japan, and Korea), Middle Asia (Iran and Turkey), and South America (Brazil and Mexico). If the subjects of a study came from two or more of the above-listed regions, the trial was classified as multinational. The baseline information from the different geographic regions is summarized in Table 2. Most AD trials were conducted in Western countries, and their demographic characteristics were comparable to those of East Asian trials. The only exception was that more female patients were included in East Asian trials. Middle Asian trials enrolled a higher proportion of male patients with AD than other regions. The baseline ADAS-cog score was highest in the South American population, while a similar distribution was found among other regions.

Detailed demographic information of the included studies is listed in Additional file Table 1.

**Disease progression model**

Most of the ADAS-cog data points included in the analysis were measured for two years, except for four observations derived from one study [30]. Only placebo data acquired during a period two years were modeled because long-term observations might have biased the estimation of disease progression rate. The results of the structural model exploration showed that the disease progression model combined with the $E_{\text{max}}$ model to describe the placebo effect was the most appropriate basic model because of its lowest AIC value and higher precision of parameter estimations compared with those using the exponential or sigmoid $E_{\text{max}}$ model. When using the inverse Bateman function, the offset rate constant of the placebo effect was difficult to estimate because the duration of some clinical trials included in this study less than 12 weeks, and the minimization process for parameter estimation failed to converge successfully.

The parameter estimates of the final model are shown in Table 3. Covariate screening analysis demonstrated that both baseline ADAS-cog score and age had a significant impact on the disease progression rate ($P < 0.05$). The relationship between the individual estimate of the disease progression
rate from the basic model and the baseline ADAS-cog score showed an obvious nonlinear correlation, as shown in Figure 2A. In this study, the inverse U-type model was superior to the power model for assessing the baseline ADAS-cog effect as the higher degree of OFV reduction compared with the basic model (10.884 versus 8.595), and the inverse U-type model was more consistent with the nonlinearity for boundary scale data, which is characterized by the faster disease progression rate in the moderate AD population than in the mild and severe AD populations. In addition, covariate analysis also found a negative correlation between age and disease progression rate (Figure 2B). Younger patients with AD deteriorated faster in cognitive function than elderly AD patients. Compared with the basic model, the inter-study variability of disease progression rate decreased from 18.5 % to 14.8 % after the introduction of the effects of ADAS-cog and age as covariates. The final mathematical expression of the population estimate of $\alpha$ is shown in Equation 7. For the AD population with an average age of 73.5 years and ADAS-cog baseline score of 24.5 points, the typical value of change rate in ADAS-cog scale was estimated to be 0.112 points/week (5.82 points/year). Theoretically, the disease progression rate was the fastest when the ADAS-cog score reached the inflection point of 35.5. The curved surface among the ADAS-cog baseline score, age, and population estimate of $\alpha$ are shown in Figure 2C.

[Please see the supplementary files section to view the equation.] (7)

In the final model, the population estimate of $\beta$ was -1.87 points. The inter-study variability of $\beta$ was estimated to be 72.5 %, indicating that the placebo effect varied largely across trials. The sign of the individual estimate $\beta$ was positive in only seven studies, which suggested that cognitive function deteriorated more in these studies and it was unclear whether this could be attributed to the nocebo effect. The estimated population value of $ET_{50}$ was 7.99 weeks and its inter-study variability was fixed at 0 because of the very large shrinkage value. Thus, it was anticipated that the plateau of the placebo effect might be reached at approximately 16 weeks in clinical trials. In addition, a significant negative correlation was found between the inter-study variability of $\alpha$ and $\beta$, and the corresponding Pearson correlation coefficient was -0.717.

It was not found that add-on design and the proportion of subjects receiving background treatment during the trial and publication year had a significant impact on the disease progression rate. In addition, we did not find any factors that had a significant effect on the theoretical maximum placebo effect. Since information regarding duration of education, NPI score, AD duration, proportion of APOE $\varepsilon$4 carriers, and proportion of whites were not reported in more than 30 % of the studies, these variables were not included in the covariate screening step.

Subgroup analyses
The observed mean change from baseline in the ADAS-cog score under different stratifying factors is shown in Figure 3. The results of the subgroup analysis of disease progression rate $\alpha$ and maximum placebo effect $\beta$ are shown in Table 4. It was revealed that the disease progression rate published before 2008 was significantly higher than that published after 2008. However, after correcting for baseline ADAS-cog score and age effect, this significant difference disappeared (Table 4). This is why the publication year could not be introduced into the model as a covariate. Because the age of subjects was comparable between the trials published before or after 2008, we hypothesized that the decline rate of ADAS-cog score in the placebo group tended to decrease in recent AD trials, which might be driven by the relatively lower baseline ADAS-cog score. In addition, no difference was found between trials using the add-on design and those not using the add-on design.

An interesting finding was that disease progression varied across geographic regions. Even after correcting for the baseline ADAS-cog score and age effect, trials conducted in Middle Asian countries had a higher disease progression rate than trials conducted in international and East Asian countries. East Asian trials showed the slowest disease progression rate compared to other trials. The theoretical maximum placebo effect in international multiple trials was significantly higher than that in North American populations, while the Middle Asian population showed the weakest placebo effect than any other region. Based on the model parameters obtained by subgroup analyses, we simulated the typical time course of change in ADAS-cog score from baseline under different stratification factors (Figure 4).

**Model evaluation and validation**

The median values of the model parameters obtained from the non-parametric bootstrap analysis were consistent with the parameter estimates of the final model. In addition, the 95 % CI of the covariate correction factors did not contain 0, further verifying the reliability of the covariate analysis results.

The quality of the fit of the final model was relatively good (Additional file Figure 2). Both population predictions and individual predictions correlated well with the observations. The model predictions were evenly distributed on both sides of the standard line without obvious deviation. The absolute values of the conditional weighted residuals were less than 6 and no systematic trend was found over time and the population predictions. The visual predictive check (Additional file Figure 3) showed that the median and the 2.5 % and 97.5 % quantiles of the observations fell within the 95 % CI of the quantile predicted by the final model, indicating that the final model could reflect the central trend and variability of the observation values. Thus, the final model had good reproducibility for the original data and good predictive capabilities.

**Discussion**

This model-based meta-analysis integrated longitudinal data of ADAS-cog in the placebo group from published AD clinical trials from 1992 to 2019, extended the observation period to two years, and
reconstructed the literature-level AD disease progression model. We found that baseline ADAS-cog score and age were two important factors influencing the disease progression rate, consistent with the results of individual-level studies based on the ADNI database [16, 31].

Previous studies based on individual data have found that the relationship between disease progression rate and baseline ADAS-cog score is nonlinear [32, 33]. Therefore, we used the inverse U-type model in this study to describe the nonlinear relationship between disease progression rate and ADAS-cog score. However, since the maximum baseline ADAS-cog score of the subjects included in this study at summary level was no more than 40 points, the trend that the disease progression rate decreases with increase in the baseline ADAS-cog score was not obvious. Thus, it is necessary to be careful to extrapolate the disease progression rate when the baseline ADAS-cog score is greater than 40 points at the summary level. Compared with the study by Ito et al., our study found that baseline age also had a significant impact on the disease progression rate. The estimated correction coefficient for the age effect was -2.17, which was slightly lower than the value of -1.8 obtained by using individual data modeling [13]. The cognitive decline in the younger AD population was faster than that of the elderly AD population. For example, the disease progression rate of the 60-year-old AD population was estimated to be about 1.87-times as fast as that of the 80-year-old AD population. Based on the sample size, baseline ADAS-cog score, and age, the final model could predict the time course of ADAS-cog in the placebo group as a historical control in future AD trials, whereas model predictions were compared with observations from clinical trials to aid in the evaluation and interpretation of trial outcomes [25, 34, 35].

The guidelines for AD published around 2008 recommend CHEIs and memantine as the standard treatment for AD patients [1, 2]. The number of AD trials designed with these drugs as add-on treatment has increased since then. In this study, we found that the disease progression rate was almost the same between trials with add-on design and non-add-on design. This is consistent with the conclusion that currently-approved drugs do not change the disease progression rate. The enrolled subjects were required to have been treated with these drugs at a stable dose for at least one to three months before entering the add-on trials. The drug efficacy was nearly close to the plateau at the beginning of the trial and it was expected not to affect the estimation of the placebo response.

In this study, subgroup analysis of the disease progression rate according to the year of publication was conducted to test the hypothesis that clinical care and patient management may improve over time. We found that the mean annual increase in ADAS-cog score in AD trials published after 2008 was significantly lower than that before 2008. However, when the baseline ADAS-cog score was corrected, the impact of the publication year on the disease progression rate disappeared, which suggested that the publication effect was due to the fact that more patients with mild AD were included in recent trials. The above-described results suggest that when performing systematic review or meta-analysis, the data of add-on trials and non-add-on trials as well as the data published in the past ten years and data published more than ten years ago can be combined and analyzed to increase the richness of the available data.
Clinical trials of AD have been conducted globally, and we found that clinical trials in Middle Asian countries showed rapid disease progression in the placebo group (Table 4). It should be noted that the sample size of trials in Middle Asia was relatively small, which might have resulted in a biased estimation of the placebo decline rate. Meanwhile, by comparing the original and adjusted disease progression rates, we concluded that the higher decline rate in South America trials was due to the inclusion of patients with more severe AD. Importantly, we found that the covariate-adjusted disease progression rate in East Asia was significantly lower than that in North America. This conclusion is consistent with a previous study [24], in which, however, the baselines of subjects in different regions were not comparable and the conclusion was questioned. Cerebrovascular insufficiency was associated with the onset and progression of cognitive dysfunction [36], and we suspected vascular load difference between western and eastern populations might contribute to the relatively slower decline rate in East Asian countries. Final model simulation suggested that the ADAS-cog change of East Asian populations was flatter than that of North American populations. The typical change scores from baseline in ADAS-cog of East Asian and North American populations at six months were 0.9 (95 % CI: 0.03–1.68) and 1.91 (95 % CI: 1.75–2.06), respectively. Therefore, differences in disease progression and placebo effects from different regions, especially in East Asia and North America, need to be noted when conducting multicenter clinical trials. However, it should be noted that most of the ADAS-cog data points in East Asia included in the analysis were measured in six months, and the trajectory of disease progression in East Asia over a longer treatment duration still requires further data support.

This study has some limitations. First, we did not examine the effects of some important covariates on disease progression rate. For example, there is substantial evidence that APOE ε4 alleles are high risk factor for AD progression [15]. Other important biomarkers, such as the ratio of p-tau$_{181}$P over Aβ$_{1–42}$ in the cerebrospinal fluid and hippocampal volume [15, 17, 18], was also not possible to account for because the proportion of studies that did not include these key covariates was too high to be integrated into the model. Second, the extrapolation of the final model to predict ADAS-cog changes after two years might be limited. There were some deviations between model predictions and observations when predicting long-term placebo response in mild-to-moderate AD subjects ((Additional file Figure 4). Finally, the time lag between recruitment of AD patients into clinical trials and the published year might interfere with drawing the conclusion of the publication year effect [37].

Conclusions

To our knowledge, this study includes the highest number of clinical trials so far in the field of AD disease progression modeling. We found that the baseline ADAS-cog score and age are two important predictors for the disease progression rate in AD. The disease progression of AD is not affected by publication year and add-on design but shows different regional differences. The disease progression rate in East Asia is significantly lower than that in North America. These conclusions should be fully considered when designing future AD trials or conducting indirect comparisons of clinical outcomes.
List Of Abbreviations

AD: Alzheimer's disease

ADAS-cog: cognitive subscale of the Alzheimer’s Disease Assessment Scale

aMCI: amnestic mild cognitive impairment

CHEIs: cholinesterase inhibitors

CI: confidence interval

MMSE: Mini-mental State Examination

NPI: neuropsychiatric inventory

VPC: visual predictive check

Declarations

Ethics approval and consent to participate

Not applicable

Consent for publication

Written informed consent for publication was obtained from all participants.

Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

This was not an industry supported study. The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding
This work was received financial support from National Major S&T Project (2018ZX09734005, 2018ZX09711001-009-011, 2018ZX09731016, 2017ZX09304003, and 2018ZX10303501), the project of Shanghai Municipal Health Planning Commission (2018YQ48), and Shanghai S&T Innovation Plan (17401970900).

Authors' contributions

Ningyuan Zhang, Lujin Li and Qingshan Zheng designed the workflow of this study. Ningyuan Zhang and Hongxia Liu searched and extracted the literature-level data and established the dataset, Qingshan Zheng was responsible for the verification of the consistence of the data extracted from the two workers. Ningyuan Zhang and Xijun Zheng analyzed and modelled the data. Ningyuan Zhang and Lujin Li were major contributors in writing the manuscript. Xijun Zheng also provided very valuable insights to the writing work. All authors read and approved the final manuscript.

Acknowledgments

The authors wish to acknowledge the participants and their study partners who volunteered their time to complete this study.

Authors' information

Qingshan Zheng, Lujin Li and HongXia Liu were professors of ShangHai University of Traditional Chinese Medicine, Ningyuan Zhang and Xijun Zheng were postgraduates.

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Tables

Table 1 Demographic characteristics of the included studies
### Table 2 Characteristics of the included studies across different regions

| Region                  | International | North America | Europe/Oceania | East Asia | Middle Asia | South America |
|-------------------------|---------------|---------------|----------------|-----------|-------------|---------------|
| Number of trials        | 56            | 56            | 15             | 5         | 5           | 3             |
| Total sample size       | 10842         | 6855          | 799            | 457       | 98          | 159           |
| Sample size per arm*    | 135 (19, 639) | 102 (11, 746) | 40 (14, 153)   | 37 (19, 268) | 20 (15, 25) | 18 (12, 129) |
| Age, year*              | 73.2 (68.3, 78.3) | 75.0 (58.0, 81.7) | 72.1 (64.6, 79.8) | 74.5 (69.0, 75.4) | 73.1 (67.7, 73.7) | 72.6 (71.7, 74.5) |
| Gender, male%*          | 40.3 (30, 59) | 45.0 (0, 98)  | 38.0 (24.0, 53.3) | 31.8 (23.8, 37.0) | 52.1 (45, 60) | 38.9 (27.1, 41.7) |
| APOE ε4 carrier, %*     | 58.0 (0, 72) | 60.0 (41.3, 100) | 51.1 (44.0, 58.8) | N.A.      | N.A.        | N.A.          |
| ADAS-cog 11, point*     | 24.7 (18.9, 35.8) | 23.4 (13.1, 32.2) | 24.5 (15.4, 34.3) | 26.9 (23.9, 33.5) | 25.6 (17.1, 33.5) | 36.7 (29.3, 39.0) |
| NPI score, point*       | 10.3 (6.2, 21.8) | 8.65 (5.0, 16.1) | 9.2 (7.1, 11.2) | 9.1 (8.8, 9.3) | N.A. | N.A. |
| MMSE score, point*      | 19.4 (15.5, 22.9) | 20.3 (17.0, 25.4) | 19.3 (16.8, 23.6) | 16.1 (14.6, 16.6) | 13.2 ** | 17.4 (17.2, 17.6) |

*reported mean data were express as median (minimum-maximum)

**only one study reported the relevant information

N.A.: no available information was reported
Table 3 Parameter estimates of the final disease progression model

| Parameters          | Estimates (RSE%) | NONMEN 95% CI | Bootstrap Median | Bootstrap 95%PI |
|---------------------|------------------|---------------|------------------|-----------------|
| **Fixed effect**    |                  |               |                  |                 |
| \( \alpha \), point/week | 0.112 (6.5)     | 0.098 ~ 0.126 | 0.114            | 0.102 ~ 0.139   |
| \( \beta \), point   | -1.87 (26.4)     | -2.84 ~ -0.902| -2.03            | -4.28 ~ -1.27   |
| \( ET_{50} \), week  | 7.99 (50.4)      | 0.091 ~ 15.9  | 8.91             | 3.22 ~ 25.4     |
| **Covariate**       |                  |               |                  |                 |
| \( \theta_{\text{baseline ADAS-cog score}} \) | 1.53 (34.8)     | 0.487 ~ 2.57  | 1.46             | 0.581 ~ 2.96    |
| \( \theta_{\text{baseline age}} \) | -2.17 (29.1)     | -3.41 ~ -0.931| -1.95            | -3.73 ~ -0.625  |
| **Random effect**   |                  |               |                  |                 |
| \( \eta_1 \) for \( \alpha \), % | 14.8 (43.0)     | 2.33 ~ 27.3   | 15.5             | 2.9 ~ 27.3      |
| \( \eta_2 \) for \( \beta \), % | 72.5 (15.2)     | 50.9 ~ 94.1   | 74.4             | 53.3 ~ 106.6    |
| Pearson correlation  | -0.717           |               | NA               |                 |
| **Residual error**  |                  |               |                  |                 |
| \( \sigma_{\text{add on, point}} \) | 5.64 (6.8)       | 4.89 ~ 6.39   | 5.54             | 4.83 ~ 6.26     |

Table 4 Subgroup analysis of disease progression rate

| Uncorrected disease | Covariate corrected disease | Maximum extent of placebo effect, point |
|---------------------|-----------------------------|----------------------------------------|
| progress rate, point/year | progress rate, point/year | Mean (95% CI) |
| Mean (95% CI)       | Mean (95% CI)              | -1.89 (-2.08, -1.69)                   |
| <2008                | 6.13 (5.89, 6.39)          | 5.92 (5.80, 6.04)                      |
| >=2008               | 5.56 (5.40, 5.73)          | 5.81 (5.72, 5.92)                      |
| **Design**           |                             |                                        |
| add on               | 5.53 (5.26, 5.80)          | 5.85 (5.69, 6.00)                      |
| on add on            | 5.84 (5.64, 6.04)          | 5.87 (5.74, 6.01)                      |
| **Region**           |                             |                                        |
| North America        | 5.81 (5.66, 5.97)          | 5.78 (5.67, 5.90)                      |
| America              | 5.62 (5.35, 5.89)          | 6.11 (6.00, 6.23)                      |
| East Asia            | 5.67 (5.15, 6.19)          | 5.45 (4.93, 5.97)                      |
| Asian Pacific        | 5.58 (4.68, 6.48)          | 5.54 (4.79, 6.29)                      |
| East Asia            | 5.67 (5.15, 6.19)          | 5.45 (4.93, 5.97)                      |
| Middle East Asia     | 7.00 (5.98, 8.02)          | 6.76 (6.20, 7.31)                      |
| America              | 7.27 (6.49, 8.06)          | 6.27 (5.48, 7.06)                      |

The values have been converted to annual disease progression rate, \( \alpha \)(point/week)\times 52 week/year
Figures

Figure 1

The flow diagram illustrating the inclusion and exclusion of studies into the final analysis.
Figure 2

(A) Plot of the individual estimate of disease progression rate $\alpha$ derived from the base model versus the baseline ADAS-cog score. The two parameter-covariate relationships, power and U-type function, were demonstrated by using blue and red lines, respectively. (B) Plot of the individual estimate of disease progression rate $\alpha$ derived from the base model versus the baseline age. The blue fitted line demonstrated the correlation between them. In plot (A) and (B), each black circle stands for one independent study and the circle size is proportional to the corresponding sample size. (C) A surface diagram to describe the relationship among age, ADAS-cog score, and the disease progression rate. Younger patients with higher baseline ADAS-cog score were predicted to manifest faster cognitive decline.
Figure 3

Observed mean change from baseline in ADAS-cog score under different stratification scenes, the point size is proportional to the corresponding sample size.: (A) published before 2008 versus after 2008, The red points and blue points stand for trials published after 2008 and before 2008, respectively. (B) add-on design versus non add-on design, the red points and blue points stand for add-on trials and trials not using add-on design respectively. (C) different geographic locations.
Figure 4

Typical time course of the change of ADAS-cog score from baseline under different stratification scenes: (A) published before 2008 versus after 2008, (B) add-on design versus non add-on design, (C) different geographic locations. The shadows are the model predicted 95% CI of the placebo response. The solid line is the typical value of the model predicted placebo response.

Supplementary Files

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