European multicentre double-blind placebo-controlled trial of Nilvadipine in mild-to-moderate Alzheimer’s disease—the substudy protocols: NILVAD frailty; NILVAD blood and genetic biomarkers; NILVAD cerebrospinal fluid biomarkers; NILVAD cerebral blood flow

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ABSTRACT

Introduction: In conjunction with the NILVAD trial, a European Multicentre Double-Blind Placebo Controlled trial of Nilvadipine in Mild-to-Moderate Alzheimer’s disease (AD), there are four NILVAD substudies in which eligible NILVAD patients are also invited to participate. The main NILVAD protocol was previously published in BMJ Open (2014). The objectives of the NILVAD substudies are to determine whether frailty, cerebrospinal fluid (CSF), blood biomarker profile and Apolipoprotein E (APOE) status predict response to Nilvadipine, and to investigate the effect of Nilvadipine on cerebral blood flow and blood biomarkers.

Methods and analysis: All participants who fulfil criteria for the main NILVAD study are eligible for participation in the NILVAD substudies. Participation is subject to informed consent and whether the substudy is available at a particular NILVAD study site. Each substudy entails extra measurements during the course of the main NILVAD study. For example, in the blood and genetic biomarkers substudy, extra blood (30 mL) will be collected at week 0, week 13, week 52 and week 78, while in the cerebral blood flow substudy, participants will receive an MRI and transcranial Doppler measurements at week 0, week 26 and week 78. In the CSF substudy, 10 mL CSF is collected at week 0 and week 78.

Ethics and dissemination: All NILVAD substudies and all subsequent amendments have received ethical approval within each participating country, according to national regulations. Each participant provides written consent to participate. All participants remain anonymised throughout and the results of each substudy will be published in an international peer reviewed journal.

Trial registration number: EUDRACT 2012-002764-27; Pre-results.

Strengths and limitations of this study

▪ These nested NILVAD substudies will collect biomarker data to determine if biomarker status predicts response to Nilvadipine.
▪ The substudies run in parallel with the main NILVAD trial and thus facilitate the collection of data.
▪ This biomarker data collection in mild-to-moderate Alzheimer’s disease (AD) will provide valuable insight into the in vivo effects of Nilvadipine on these parameters. In addition, these studies provide the opportunity to study the relationship between the different outcome measures.
▪ The substudies will not necessarily be completed by all participants recruited for the main NILVAD study (500 participants). Therefore, the results are potentially less generalisable compared to the main NILVAD study. For example, the cerebral blood flow substudy will only run in one study site and aims to recruit 40 patients.

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INTRODUCTION

The NILVAD study is a large European multicentre, randomised, double-blind, placebo-controlled trial investigating the efficacy and safety of Nilvadipine as a disease course modifying treatment in mild to moderate Alzheimer’s disease (AD). As part of this trial, four nested substudies have been developed to investigate if frailty status, cerebrospinal fluid (CSF) and blood biomarker profiles, and Apolipoprotein (APOE) status can predict response to Nilvadipine. Furthermore, the in vivo effects of Nilvadipine on frailty, cerebral blood flow and blood biomarker status over time will be determined. The substudies have also been designed to examine other end points that are relevant to AD pathology and allow assessment of AD progression. The four substudies are detailed as follows.

Frailty substudy

Frailty is a common and important syndrome that is increasingly prevalent with advancing age. A consensus view is emerging where the physical phenotype of frailty develops as a consequence of complex biological interactions that promote cell senescence, leading to a cumulative decline in multiple physiological systems, particularly the brain, endocrine system, immune systems and skeletal muscles. Furthermore, there is accumulating evidence to support a temporal association between frailty, cognitive impairment and dementia.1 2 It has been suggested that frailty can be used as an outcome measure because of its relationship with survival and adverse outcomes such as hospitalisation in older adults.3

Thus, a substudy to measure frailty was developed aiming (1) to investigate whether baseline frailty status predicts response to treatment with Nilvadipine and (2) to determine if treatment with Nilvadipine influences frailty status. Frailty will be measured via gait speed and a frailty index.

A large study by Fried et al4 investigating frailty in older adults, operationalised frailty using a cluster of variables: unintentional weight loss, self-reported exhaustion, low energy expenditure, weak grip strength and slow gait speed. Their study provided the basis for identifying frailty phenotypes to predict outcome. Rockwood et al5 found that slow gait speed had the strongest association with dependency, long-term care and death, indicating gait speed might be a useful single indicator of frailty. Therefore, gait speed was chosen as a prospective physical marker of frailty in this substudy.

The Rockwood frailty index (FI) was developed from part of the Canadian Study of Health and Ageing.5 It consists of multiple parameters (‘deficits’) that can be measured, and allows for calculation of the presence or absence of frailty (simple calculation of the presence or absence of each variable as a proportion of the total: for example, 20 deficits present out of a possible 70 gives a FI of 20/70=0.29). Older adults with a FI of ≤0.08 can be considered non-frail, whereas those with an FI of ≥0.25 are considered frail. Considering that parameters for the FI should be biologically sensible, accumulate with age and do not saturate too early, a list of parameters will be collected to define a valid FI before, during and after participation in the NILVAD study.6 7 They are extracted from existing questionnaires, including the Disability Assessment of Dementia (DAD), Alzheimer’s Disease Assessment Scale (ADAS-Cog 12), the Lubben Social Network Scale (LSNS) and the Charlson Comorbidity Index (CCI).8–11

We expect that if Nilvadipine has a positive effect on cognition, it will also do so on frailty, with less deterioration or with possible stabilisation.

Blood and genetic biomarkers substudy

Nilvadipine has been shown to have a multimodal action, decreasing amyloid production in transgenic mouse models of AD and increasing clearance of amyloid across the blood-brain barrier (BBB) as well as having anti-inflammatory effects in in vitro models.12–14 Thus, changes in amyloid and τ fragments along with other inflammatory markers in the peripheral blood from study baseline to end point may be useful in improving our understanding of the biological effects of Nilvadipine treatment in AD.

Levels of plasma Abeta (Aβ) and τ have been suggested as possible biomarkers for detecting conversion from normal cognition to mild cognitive impairment (MCI) or AD.15 16 As such, measurement of plasma Aβ and τ could be useful as potential biomarkers for detecting disease-modifying therapeutic effects.

The APOE ε4 allele is one of the major genetic risk factors for AD and occurs in nearly 60% of all AD cases.17 Most studies investigating treatment strategies for AD now include APOE testing to determine whether treatment outcomes vary by genotype.18 In a study by Kennelly et al,19 AD indicated that non-APOE ε4 carriers showed more of a short-term benefit from Nilvadipine treatment than did ε4 carriers.

Recently, a broad lipidomic characterisation of other phospholipids (PL) showed that a ratio of arachidonic acid to docosahexaenoic acid (DHA) was increased within several PL classes in APOE ε4 carriers several years prior to diagnosis of MCI/AD.20 As such, we expect that identification of APOE status and monitoring of these PL species could assist in monitoring effects of Nilvadipine on AD progression.

Inflammation is an active component of AD pathology, with aberrant activation of microglia and astroglia resulting in prolonged exposure to proinflammatory cytokines, contributing to chronic neuroinflammation in the brains of patients with AD. Studies have shown that various proinflammatory cytokines are increased in plasma and CSF in AD compared to in control subjects.21 22 Preclinical work on Nilvadipine supports its potential anti-inflammatory action and therefore examination of proinflammatory cytokines may be useful.
biomarkers in determining therapeutic response to Nilvadipine.

The aim of this substudy is thus to determine whether genetic influences can modify the effect of Nilvadipine on AD pathogenesis and whether changes in plasma Aβ, τ and inflammatory markers from study baseline to end point are altered by Nilvadipine treatment.

**CSF substudy**

CSF biomarker levels of total τ (T), phosphorylated τ (P) and β-amyloid 1-42 (Aβ1-42), reflect, respectively, axonal nerve cell degeneration, formation of neurofibrillary tangles and disturbed β-amyloid metabolism, thus allowing identification of neurodegenerative and cerebrovascular disease mechanism. These biomarkers, T, P, τ, Aβ and the Aβ/τ ratio, are considered to be most closely associated with AD core pathology. A systematic review of CSF biomarkers in AD, in 2003, demonstrated a diagnostic specificity of ~90% and a sensitivity of ~80% for both T and P for AD.23 In addition, it has been shown that the Aβ/τ ratio can be used to predict the conversion of MCI to AD with a specificity of 85% and a sensitivity of 86%.24 Although pure AD is often regarded as the most common dementia, it is probably less common than mixed dementia (MD: presence of AD and small vessel disease).25 A different biomarker pattern is characteristic of patients with manifest MD as well as pure subcortical vascular dementia with increased blood flow measures in AD. The substudies will also determine whether biomarker and genetic status predicts response to Nilvadipine, and will provide clues to the mechanism of action of Nilvadipine in AD. In addition, NILV substudies will provide novel data on the effects of Nilvadipine on frailty. Finally, the long-term storage and retention of blood and CSF samples will contribute significantly to a unique bioresource for studies of AD in the future.

In conclusion, determination of CSF biomarkers may be useful for enhanced diagnostic purposes to assess which patients display signs of pure AD, mixed type dementia or a more pure vascular profile, with expectation that the treatment response is larger in patients with vascular involvement.

Thus the aim of this CSF substudy is to explore the biomarker status of NILVAD participants at baseline as a predictor of response to Nilvadipine and investigate biomarker changes over time, with Nilvadipine treatment, possibly enhancing our understanding of the in vivo effect of Nilvadipine.

**Cerebral blood flow study**

Animal models indicate that AD may be associated with profound vascular changes.29 For example, cerebral autoregulation—the mechanism that aims to stabilise cerebral blood flow during changes in blood pressure—was severely impaired in the transgenic mouse model for AD.30 Specifically, Aβ may lead to both, structural and functional cerebrovascular changes that may result in an increased risk for cerebral hypoperfusion and ischaemia. A recent study found that Nilvadipine inhibited vasoconstriction elicited by Aβ and restored decreased cortical perfusion in a transgenic mouse model, overexpressing human Aβ.31 How AD affects the cerebral vasculature and cerebral autoregulation in humans remains poorly understood. Nilvadipine has been shown to significantly improve memory and regional cerebral blood flow (rCBF) deficits in the left frontal lobe in hypertensive patients with MCI, while amlodipine did not.31 In fact, regional cerebral blood flow in the temporal lobe of the amlodipine group deteriorated. The rCBF and cognitive effects were independent of the reduction in systolic and diastolic blood pressure after treatment.

In addition to the potentially beneficial cerebrovascular effects of Nilvadipine, it may also improve blood pressure (BP), with variation in systolic BP shown to be reduced by calcium-channel blockers (and non-loop diuretic drugs), but increased by ACE inhibitors, angiotensin-receptor blockers and β-blockers.32 Thus, BP variability, a factor that may enhance risk of cerebral ischaemia, could be reduced by Nilvadipine, more than by other antihypertensive drugs.

Thus this CBF substudy will investigate the effects of Nilvadipine on CBF, BP and BP variability in study participants. It is expected that Nilvadipine treatment, through direct systemic and cerebral vascular effects, will beneficially modulate these characteristics.

In summary, these substudies will provide information on the effects of Nilvadipine on physical, cardiovascular and blood flow measures in AD. The substudies will also determine whether biomarker and genetic status predicts response to Nilvadipine, and will provide clues to the mechanism of action of Nilvadipine in AD. In addition, NILV substudies will provide novel data on the effects of Nilvadipine on frailty. Finally, the long-term storage and retention of blood and CSF samples will contribute significantly to a unique bioresource for studies of AD in the future.

**METHODS AND ANALYSIS**

**Frailty substudy**

**Study design**

Study participants will undergo a frailty assessment at particular time points. The schedule consists of completion of the DAD, ADAS-Cog12, LSNS and CCI. Collection of data variables from these assessments will facilitate definition of a FI before, during and after participation in the NILVAD study.

The main NILVAD study runs for a total of 82 weeks, with 78 weeks of study drug treatment (Nilvadipine or placebo). The LSNS and CCI (figure 1) are performed at study baseline (week 0) and treatment end point (week 78). The DAD and ADAS-Cog and gait speed (figure 2) are performed at week 0, 13, 52 and 78.

**Inclusion/exclusion criteria**

The inclusion and exclusion criteria for this substudy are as for the main NILVAD study.33
<table>
<thead>
<tr>
<th>Frailty Assessment Time points</th>
<th>Visit 2</th>
<th>Visit 4</th>
<th>Visit 7</th>
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<td>Week 13</td>
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<tr>
<td>Charston Comorbidity Index</td>
<td>Assessment of what additional disorders the patient may have</td>
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<td>ADAS Cog</td>
<td>Items 9, 10 and 11</td>
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**Figure 1** Frailty assessment schedule. ADAS, Alzheimer’s Disease Assessment Scale; DAD, Disability Assessment of Dementia.

**Figure 2** Protocol for measuring gait speed.

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**Preparations**
Make 4 markings on the ground with tape (indicated above in red. Choose a suitable hallway, where you can do all the measurements):

**Outcome**
Walking speed over 4 meters (between marking 2 and 3)

**Who**
Trained research personnel

**Material**
- Stopwatch with 2 decimals precision
- Paper
- Pen

**Method**
Inform the patient that you are going to measure his/her normal walking speed. To do so, the patient needs to walk from marker 1 to 4. This way, you will eliminate starting/stopping from the measurement.

1. Ask the patient to stand at marker 1. Indicate the patient to start walking.
2. As soon as the first foot touches or crosses marker 2, you start the stopwatch.
3. As soon as the last foot has crossed marker 3, you stop the stopwatch.
4. Tell the patient to stop when he/she reaches marker 4.
5. Indicate the time (with 2 decimals) and the walking aid.

Patients should be measured as they normally walk. If this is with a walking aid, then measure them with the walking aid. Indicate with the time, which kind of walking aid they used.
Recruitment
There are a total of 23 study sites in 9 participating European countries. Recruitment for the substudies will be undertaken locally at each study site according to the local guidelines and procedures relevant to that site. All study sites will participate in the Frailty substudy, excluding Germany.

Consenting process
All consenting participants for the main NILVAD study will be invited to also provide written consent to partake in the substudies, with the option of consenting to participate in as many of the substudies as they wish, dependent on availability of the substudy at any given study site. Consent will be obtained following a full explanation of each substudy, its objectives, risks and potential benefits. A patient information leaflet and a carer’s information leaflet will be provided at least 7 days prior to obtaining written consent, to allow participants and their carers time to consider enrolling. Written consent will be obtained at the main NILVAD screening visit, prior to the undertaking of any interventions or assessments.

As per the consenting process for NILVAD, when a participant has reduced decision-making capacity, the procedure for obtaining consent must follow national law and will be assessed by the relevant bodies in each of the participating countries.

Sample size calculation
The FI ranges from 0 to 1, with a natural maximum of 0.7. To calculate the sample size, we conservatively estimated the variance to be large, that is, a SD of 0.3. A sample size of 140 is adequate to allow detection of a 10% difference in FI between the NILVAD (experimental) and placebo (control) groups, and facilitate rejection of the null hypothesis that the population means of each group are equal with probability (power) 0.8. The type I error probability associated with the test of this null hypothesis is a two-sided $\alpha=0.05$.

Data analysis
Data will be automatically drawn from an online Macro system, using a script in Matlab (The Mathworks, Natick, Massachusetts, USA). Then the FI will be calculated as a weighted score based on items from the ADAS-Cog (3 items), CCI (15 items), DAD (36 items), LSNS (2 items) and the gait speed (1 item). Points scored per visit are divided by the number of items applied at that visit, resulting in a FI between 0 and 1.

First, an independent samples t-test will be used to assess if there is a difference in FI between the groups at baseline, using $\alpha=0.05$ as significance level. When the scores do not differ significantly, an independent samples t-test will be used to assess if there is a difference in FI between the groups at week 78, using $\alpha=0.05$ as significance level. If the FI scores do differ at baseline between the groups, the data will be analysed using a mixed linear model (MLM), to examine whether Nilvadipine treatment (independent fixed factor) altered repeat measurements of FI over time (outcome factor) while controlling for random effects (each participant), again using $\alpha=0.05$ as significance level.

The above analyses will be conducted on the ‘per protocol population’ (those participants who have no violations to the protocol for the duration of the study).

Blood and genetic biomarkers substudy
Study design
This substudy will involve collection of 30 mL of blood for genetic and biomarker analysis at four time points: week 0, week 13, week 52 and week 78. Whole blood (1 mL) will be aliquoted for genotyping for APOE 4 at week 0. The remaining 29 mL of the sample will be processed for biomarker analysis as per the protocol standard operating procedure (SOP), for blood processing and sample preparation (box 1).

All samples will be collected, aliquoted and stored locally at each study site initially and then transported for final study analysis to a central study site, before transfer to a biobank for long-term storage. All samples will be stored under specific conditions and coded for identification according to the protocol SOP.

Inclusion/exclusion criteria
The inclusion and exclusion criteria for this substudy are as for the main NILVAD study.33

Recruitment
All study sites will participate in the blood and biomarker substudy except for those based in the UK and Germany.

Sample size calculations
Using the available serum data on decreases of Aβ40 observed in the phase I/II Nilvadipine study (unpublished data), we calculated an effect-size of 0.08 (based on the mean change from baseline to week 6 of the study, and the variance of Aβ40 measurements within treated and untreated participants). Given this effect size, a sample size of 254 participants is proposed to allow incorporation of 4 repeated samplings, to detect statistical significance at $\alpha=0.05$ with a power of 90%.

Sample analysis
Pure Gene Kits (Gentra Systems, Minneapolis, Minnesota, USA) will be used for extracting DNA from whole blood and PCR analyses for APOE will be performed using previously established methods.34 Blood Aβ40 levels will be analysed as previously described, using commercially available ELISA kits.35 Lipidomics analysis on plasma lipid extracts will be performed using LC/MS analyses previously described.34 36 Serum levels of proinflammatory cytokines, such as tumour necrosis factor-α and IL-1β, will be examined using high-sensitivity ELISA kits as per manufacturer’s instructions.
## Sample collection

For each participant, at week 0, week 13, week 52 and week 78 visit, draw a minimum of: 2×10 mL EDTA tubes for plasma.

- Use a ×10 mL serum separator tube (with silica plug) for serum.
- Obtain blood samples via standard phlebotomy procedures. Please follow the guidelines from your facility for performing these procedures.
- Draw the correct volume of blood allowing the vacuum in the tube to be exhausted.
- Gently invert each tube to mix immediately following blood collection.
- Process blood samples for genotyping, plasma and serum immediately on collection.
- It is vital that blood samples are processed immediately as the biomarker proteins start to degrade if they are left for any length of time before processing.

### Whole blood for genotyping (only to be carried out at week 0)

- Label one 1.5 mL polypropylene cryovial per patient, with a biobanking label. This label should contain the Biobank ID, the visit number, the sample type and patient PIN.
- Remove 1 mL of whole blood from one of the 10 mL EDTA tubes that were collected for plasma.
- Transfer the 1 mL of whole blood into the 1.5 mL preflabelled polypropylene cryovial.
- Immediately store samples at −80°C.
- Fill out the electronic case report form to record the date the sample was taken and the Biobank ID numbers for each of the samples.

### Plasma preparation: 2×10 mL EDTA tubes

- As the serum tubes require 20–30 min standing time before processing, begin with the plasma processing.
- Balance 2×10 mL EDTA tubes in the centrifuge and spin for 5 min at 1380 relative centrifugal force.
- Remove tubes from centrifuge and place on ice without shaking/inverting the tubes.
- Using the pipette, draw off the supernatant (upper layer) taking care not to disturb the interface, and transfer to a 15 mL polypropylene tube on ice.
- Transfer the 1 mL aliquots of plasma into PRELABELED 1.5 mL polypropylene cryovials (anticipating 10 vials to be collected).
- Freeze the 10 cryovials of plasma at −80°C. Do not snap freeze (ie, do not dip in liquid nitrogen to freeze).
- The time taken from blood draw to freezing down the plasma samples at −80°C should not exceed 15 min.
- Fill out the electronic case report form to record the date the sample was taken and the Biobank ID numbers for each of the aliquots.

### Serum preparation: 1×10 mL serum separator tube

- Let the serum 10 mL serum separator tube (with silica plug) stand at room temperature for 20–30 min to clot.
- Balance the tube in a refrigerated centrifuge and spin for 5 min at 1380 relative centrifugal force.
- Remove tube from centrifuge and place on ice without shaking/inverting the tube.
- Using a pipette, draw off the supernatant (upper layer) taking care not to disturb the interface and transfer to a 15 mL polypropylene tube on ice.
- Transfer the 1 mL aliquots of serum into PRELABELED 1.5 mL polypropylene cryovials (anticipating 5 vials to be collected).
- Freeze the 5 cryovials of serum at −80°C. Do not snap freeze (ie, do not dip in liquid nitrogen to freeze).
- The time taken from blood draw to placing the serum samples in the −80°C freezer should not exceed 35–45 min. This includes the 20–30 min of standing time required for the sample to clot.
- Fill out the electronic case report form to record the date the sample was taken and the Biobank ID numbers for each of the aliquots.

### Number of sample aliquots:

We shall have the following numbers of aliquots per patient at the following visits:

- Visit 2 (week 0):
  - 1.5 mL cryovial containing 1 mL of whole blood for genotyping.
  - 1.5 mL cryovials each containing 1 mL of plasma.
  - 1.5 mL cryovials each containing 1 mL of serum.
- Visit 4 (week 13), 7 (week 52) and 9 (week 78):
  - 1.5 mL cryovials each containing 1 mL of plasma.
  - 1.5 mL cryovials each containing 1 mL of serum.

## Data analysis

A MLM will also be employed to examine whether Nilvadipine treatment (independent fixed factor) altered repeat measurements of blood biomarkers over time (outcome factor) while controlling for random effects (each participant). MLM-based regression analysis is generally considered advantageous over other analysis of variance ANOVA due to its flexibility to accommodate fixed and random effects of the independent variables as well as incorporate dichotomous, continuous and categorical variables. For lipidomics analyses, principal component analysis (PCA) followed by MLM regression will be performed as described previously. Briefly, dimension reduction and multicolinearity issues will be resolved using PCA.

In order to ensure sampling adequacy for PCA, analysis achieving Kaiser–Mayer–Olkin (KMO) value of >0.6 and Bartlett p value <0.05 will be further investigated. Variables with eigenvalues of >1 will be retained, PCA will be used for extracting components and varimax with...
Kaiser normalisation will be used for rotation to simplify and clarify the data structure. The Anderson-Rubin method will be used for exporting uncorrelated scores in order to perform MLM regression analyses, as for other blood biomarkers.

The above analyses will be conducted on the ‘per protocol population’.

**CSF substudy**

**Study design**

The CSF substudy will involve collection of CSF via lumbar puncture (LP), for biomarker and inflammatory marker analysis (box 2). Patients will undergo a LP prior to start of the NILVAD study drug (ie, between the screening and baseline assessment (week 0), and again after treatment determination (week 78).

A paired serum albumin sample will also be drawn to allow for calculation of the albumin CSF-serum ratio (box 3). The ratio of albumin in CSF to serum is an indicator of BBB integrity. An increased ratio can indicate increased permeability and it has been demonstrated that AD pathology can lead to BBB dysfunction. While the underlying pathological mechanisms have yet to be fully elucidated, it is thought that alterations to the BBB affects Aβ clearance and may contribute to neuroinflammation seen in AD as a result of fluctuations in homeostasis.

Samples will be collected, aliquoted and stored locally at each study site initially, and then transported for final analysis to a central study site, before transfer to a biobank for long-term storage. All samples will be stored under specific conditions and coded for identification according to the protocol. An SOP will be adhered to for collection and processing of the CSF sample and the blood sample.

**Inclusion/exclusion criteria**

For this substudy, additional specific exclusion criteria regarding contraindications to LP must be adhered to, including (1) possible raised intracranial pressure, (2) bleeding diathesis, including thrombocytopenia or treatment with anticoagulant therapy.

**Recruitment**

All study sites, with the exclusion of those based in the UK, Germany and Italy, will participate in the CSF substudy.

**Consenting process**

In the case of the CSF substudy, consent by proxy is not applicable and all participants must retain the capacity to consent to undergo an LP. Of note, patients must retain the capacity to consent to undergo a second LP at the study end point.

**Sample size calculations**

Observed changes in CSF Aβ42 over a 78-week period from a published study were used to calculate power for

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**Box 2 Standard operating procedure for lumbar puncture (cerebrospinal fluid (CSF) collection and processing)**

**CSF sample collection and processing**

The lumbar puncture will be carried out BETWEEN THE SCREENING AND BASELINE VISIT (21-day window) and TREATMENT TERMINATION, BETWEEN WEEK 78 AND WEEK 82. The lumbar puncture should be performed ideally between 8:00 and 11:00.

- Local guidelines/routines for antiseptic cleansing and anaesthesia are to be followed.
- Position the patient in a reclining or sitting position with back arched.
- Use a Spinal Needle Quincke Type Point 0.7×75 mm (75–90 mm) for the LP.
- Insert the needle through the L3/L4 or L4/L5 interspace.
- In case of bleeding at the puncture site, the first 1 mL of CSF should be discarded and this should be recorded.
- Collect 10 mL of CSF from each participant. For CSF collection, use a POLYPROPYLENE TUBE ONLY (ie, 15 mL polypropylene tube).
- MIX the CSF gently by turning the tube upside down 3–4 times (cap on).
- Centrifuge the CSF in the polypropylene tube at 2000 g for 10 min at +4°C, to eliminate cells and debris.
- Transfer the centrifuged CSF into another sterile polypropylene tube, being careful to avoid disturbing the pellet of cells and debris at the end of the tube.
- Aliquot CSF in 1 mL aliquots into sterile 1.5 mL PREFERENCEAL POLYPROPYLENE CRYOVIALS with a screw cap.
- Place the tubes directly in a −80°C freezer. Do not snap freeze (ie, do not dip in liquid nitrogen to freeze).
- Fill out the electronic case report form to record the date the sample was taken and the Biobank ID numbers for each of the aliquots.
- The patient should be allowed to rest for 30 min to 1 hour after the lumbar puncture procedure.

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**Box 3 Standard operating procedure for paired serum albumin sample collection and processing**

**Serum preparation**

- Collect 5 mL of blood in a 10 mL serum separator tube (with silica plug) and allow to stand at room temperature for 20–30 min to clot.
- Balance the tube in a refrigerated centrifuge and spin for 5 min at 1380 relative centrifugal force.
- Remove tube from the centrifuge and place on ice without shaking/inverting the tube.
- Using a pipette, draw off the supernatant (upper layer) taking care not to disturb the interface, and transfer to a 10 mL polypropylene tube on ice.
- Transfer 1 mL aliquots of serum into PREFERENCEAL 1.5 mL polypropylene cryovials.
- Freeze the 5 cryovials of serum at −80°C. Do not snap freeze (ie, do not dip in liquid nitrogen to freeze).
- Fill out the electronic case report form to record the date the sample was taken and the Biobank ID numbers for each of the aliquots.
detecting potential decreases in CSF Aβ42 following Nilvadipine treatment. The effect size calculations were based on mean Aβ42 changes and associated variance observed by Salloway and colleagues. The resulting effect size was 0.67, which required a sample size of 98 for detecting statistically significant Aβ42 change from baseline to week 78, with a power of 90% at α=0.05.

Data analysis
Analysis of CSF will be performed according to standardised protocols. Aβ40, Aβ42, T and P 181 will be measured using a commercially available sandwich ELISA (INNOTEST; Fujirebio Europe NV, Ghent, Belgium) according to the manufacturer’s instructions.

The ratio between Aβ42 and Aβ40 for each patient will be calculated as the concentration of Aβ42 divided by the concentration of Aβ40 and multiplied by 100.41

First, an independent samples t-test will be used to assess if there is a difference in Aβ-ratio between the two groups at baseline, using α=0.05 as significance level. When the scores do not differ significantly, an independent samples t-test will be used to assess if there is a difference in Aβ-ratio change between the two groups at week 78, using α=0.05 as significance level.

The above analyses will be conducted on the ‘per protocol population’.

Cerebral blood flow substudy
Study design
Participants will undergo additional measurements of blood pressure (BP), cerebral autoregulation (CA) and cerebral blood flow (CBF) via MRI at baseline, week 26 and week 72.

Participants will be provided with a home blood pressure monitor (HBPM, Microlife Watch BP Home) at the screening visit and requested to take their own BP readings twice daily (morning and evening), after a 5 min period of relaxation, the week prior to their assessment visit at baseline, week 26 and week 78, for assessment of BP variability. Participants will be blinded to the BP readings (no result visible on monitor screen) and measurements will be conducted according to the European Society of Hypertension guidelines.42 Participants will be requested to bring the HBPM with them to the baseline visit (week 0) and week 26 and 78, to facilitate downloading of the BP data.

Additionally, at these visits, each participant will undergo an MRI scan for measurement of CBF, using arterial spin labelling (ASL) with multiple inflow times (10 time points). This sequence allows for the reconstruction of the blood flow over time for each voxel. A specific scanning protocol will be adhered to according to the study protocol (box 4), including additional sequences to measure, for instance, white matter integrity and presence of microbleeds. Subsequently, all participants will undergo CA measurements, again according to the study protocol (box 5), measuring CBF velocity in the middle cerebral artery, using transcranial Doppler. In addition, near infrared spectroscopy (NIRS) will be applied. During the measurement protocol, BP will be measured continuously using servo-controlled finger photoplethysmography (Finapres) and heartbeat will be measured using ECG.

All measurements will be taken in the morning to reduce the effect of diurnal variation in blood flow. Also, participants will have to refrain from intake of caffeine and alcohol, substances that can influence the blood flow measurement. Haematocrit level of every participant will be measured on the morning of the scan, since blood viscosity can influence the ASL measurement. This will not reduce intraindividual variation.

Inclusion/exclusion criteria
Specific contraindications to MRI must be adhered to, including (1) metal implants in situ and/or (2) suffering from claustrophobia. Patients who do not fit the MRI criteria can still participate in the CBF substudy, but without the MRI.

Recruitment
This substudy will only be undertaken in one study site, Nijmegen, in the Netherlands, to eliminate the variation
introduced by the use of different MRI scanners. Also, eligible patients from another site (Arnhem, the Netherlands) can be included in the substudy. These patients will not be included in the separate randomisation process.

Sample size calculation
The total number of participants is based on the power calculation computed on the primary efficacy parameter, CBF measured with ASL, as follows:

The primary variable of the study is a continuous variable, the CBF, from independent control and experimental subjects, with 50–50 distribution between the two groups. There have been very few studies using CBF as an outcome measure in an randomised controlled trial (RCT), but a previous study where two groups were compared detected a 25% difference. A recent RCT (N=27 vs N=31) detected a 36% change in CBF, using pulsed ASL at 3 T. To detect a 15% difference in (regional) CBF (using pulsed ASL at 3 T) between two groups, a group size of 20 per group should be sufficient, to be able to reject the null hypothesis that the population means of each group are equal with probability (power) 0.8. The type I error probability associated with the test of this null hypothesis is a two-sided 0.05.

Data analysis
All analyses will be performed blind to participant group.

Analysis of MRI data
Structural MRI scans will be processed using BET and FAST, both packages of the publicly available image processing software FSL. To identify grey matter (GM) and white matter (WM), and CSF components on the image, total brain volume (TBV) will be calculated as the sum of GM and WM volume (in mL), and intracranial volume (ICV) as the sum of GM and WM plus CSF. The TBV/ICV ratio will then be calculated as an estimate of brain atrophy.

CBF will be calculated for the whole brain GM. The data will be analysed using MLM, to test if the change in whole brain GM CBF (maximum flow) from baseline to week 26 is different for the two intervention groups, using α=0.05 as significance level.

Brain atrophy and blood viscosity will be added as additional model parameters. Using the multiple inflow times, per voxel, the maximum flow is calculated using BASIL, another FSL package. To avoid brain atrophy causing artificially low CBF values, partial volume correction will be used.

The above analyses will be conducted on the ‘per protocol population’.

ETHICS AND DISSEMINATION
All four NILVAD Substudies will be conducted according to Good Clinical Practice guidelines and in accordance with the Declaration of Helsinki. All substudies have been submitted for approval to the relevant independent Ethics Committees in each participating country, and have also been reviewed and will be continuously monitored by the NILVAD Ethics Advisory Board, for the duration of the main NILVAD study and substudies.

It is intended that all positive, neutral or negative results for the substudies will be published in international peer-reviewed journals. Authorship and publication will be as per the agreed NILVAD study publication policy.

SUBSTUDY STATUS
Enrolment for all substudies began with start of the main NILVAD trial, in May 2013, and the recruitment period continued until March 2015.

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