

Cerebrovasc Dis Extra 2017;7:140–152

DOI: 10.1159/000480738 Received: November 10, 2016 Accepted: August 18, 2017 Published online: October 10, 2017

© 2017 The Author(s) Published by S. Karger AG, Basel www.karger.com/cee



This article is licensed under the Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License (CC BY-NC-ND) (http://www.karger.com/Services/OpenAccessLicense). Usage and distribution for commercial purposes as well as any distribution of modified material requires written permission.

Original Paper

The Missing Link in the Pathophysiology of Vascular Cognitive Impairment: Design of the Heart-Brain Study

Astrid M. Hooghiemstra^a Anne Suzanne Bertens^{b, c} Anna E. Leeuwis^a Esther E. Bron^d Michiel L. Bots^e Hans-Peter Brunner-La Rocca^f Anton J.M. de Craen^c Rob J. van der Geest^g Jacoba P. Greving^e L. Jaap Kappelle^h Wiro J. Niessen^{d, i} Robert J. van Oostenbrugge^j Matthias J.P. van Osch^k Albert de Roos^b Albert C. van Rossum¹ Geert Jan Biessels^h Mark A. van Buchem^b Mat J.A.P. Daemen^m Wiesje M. van der Flier^{a, n} on behalf of the Heart-Brain Connection Consortium

^aAlzheimer Center & Department of Neurology, Amsterdam Neuroscience, VU University Medical Center, Amsterdam, The Netherlands; ^bDepartment of Radiology, Leiden University Medical Center, Leiden, The Netherlands; ^cDepartment of Gerontology and Geriatrics, Leiden University Medical Center, Leiden, The Netherlands; ^dBiomedical Imaging Group Rotterdam, Departments of Medical Informatics and Radiology & Nuclear Medicine, Erasmus MC, Rotterdam, The Netherlands; ^eJulius Center for Health Sciences and Primary Care, University Medical Center Utrecht, Utrecht, The Netherlands; ^fDepartment of Cardiology, Maastricht University Medical Center, Maastricht, The Netherlands; ⁹Division of Image Processing, Department of Radiology, Leiden University Medical Center, Leiden, The Netherlands; ^hDepartment of Neurology, Brain Center Rudolf Magnus, University Medical Center Utrecht, Utrecht, The Netherlands; ⁱImaging Physics, Applied Sciences, Delft University of Technology, Delft, The Netherlands; ^jDepartment of Neurology, Maastricht University Medical Center, Maastricht, The Netherlands; ^kC.J. Gorter Center for High Field MRI, Department of Radiology, Leiden University Medical Center, Leiden, The Netherlands; ¹Department of Cardiology, VU University Medical Center, Amsterdam, The Netherlands; ^mDepartment of Pathology, Academic Medical Center, Amsterdam, The Netherlands; ⁿDepartment of Epidemiology, VU University Medical Center, Amsterdam, The Netherlands

Keywords

Cerebral hypoperfusion \cdot Cardiovascular dysfunction \cdot Cognitive decline \cdot Heart failure \cdot Carotid occlusive disease \cdot Cerebral blood flow \cdot Small vessel disease

Abstract

Background: Hemodynamic balance in the heart-brain axis is increasingly recognized as a crucial factor in maintaining functional and structural integrity of the brain and thereby cognitive functioning. Patients with heart failure (HF), carotid occlusive disease (COD), and vascular cognitive impairment (VCI) present themselves with complaints attributed to specific parts

A.M. Hooghiemstra VU University Medical Center, Alzheimer Center PO Box 7057, NL–1007 MB Amsterdam (The Netherlands) E-Mail a.hooghiemstra@vumc.nl





Cerebrovasc Dis Extra 2017;7:14	10–152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

of the heart-brain axis, but hemodynamic changes often go beyond the part of the axis for which they primarily seek medical advice. The Heart-Brain Study hypothesizes that the hemodynamic status of the heart and the brain is an important but underestimated cause of VCI. We investigate this by studying to what extent hemodynamic changes contribute to VCI and what the mechanisms involved are. Here, we provide an overview of the design and protocol. **Methods:** The Heart-Brain Study is a multicenter cohort study with a follow-up measurement after 2 years among 645 participants (175 VCI, 175 COD, 175 HF, and 120 controls). Enrollment criteria are the following: 1 of the 3 diseases diagnosed according to current guidelines, age \geq 50 years, no magnetic resonance contraindications, ability to undergo cognitive testing, and independence in daily life. A core clinical dataset is collected including sociodemographic factors, cardiovascular risk factors, detailed neurologic, cardiac, and medical history, medication, and a physical examination. In addition, we perform standardized neuropsychological testing, cardiac, vascular and brain MRI, and blood sampling. In subsets of participants we assess Alzheimer biomarkers in cerebrospinal fluid, and assess echocardiography and 24-hour blood pressure monitoring. Follow-up measurements after 2 years include neuropsychological testing, brain MRI, and blood samples for all participants. We use centralized state-of-the-art storage platforms for clinical and imaging data. Imaging data are processed centrally with automated standardized pipelines. Results and Conclusions: The Heart-Brain Study investigates relationships between (cardio-)vascular factors, the hemodynamic status of the heart and the brain, and cognitive impairment. By studying the complete heart-brain axis in patient groups that represent components of this axis, we have the opportunity to assess a combination of clinical and subclinical manifestations of disorders of the heart, vascular system and brain, with hemodynamic status as a possible binding factor. © 2017 The Author(s)

Published by S. Karger AG, Basel

Introduction

With the rapid aging of the population, the prevalence of cognitive decline and dementia increases [1-4]. Although Alzheimer disease is the most common cause of dementia, vascular disease is increasingly recognized as an independent contributor to cognitive impairment [1]. The term vascular cognitive impairment (VCI) has been introduced to describe the complete spectrum of cognitive disorders (mild and major) associated with and due to cerebrovascular disease [1, 5–8]. VCI can be the result of irreversible structural damage to the vascular system in the brain [9]. Based on that view, treatment for VCI is often restricted to secondary prevention by treating risk factors, such as high blood pressure [10]. Recent findings suggest that cerebral hypoperfusion can also hinder the function of the brain before structural damage occurs [9]. The latter is supported by the finding that nondemented patients with cardiovascular disease show cognitive decline [11] and that in patients with heart failure (HF) cognitive functioning can be enhanced by improving cardiac function [12-14]. Furthermore, the observation of cognitive impairment in carotid occlusive disease (COD) suggests a relationship between reduced cerebral blood flow (CBF) and cognitive functioning [15]. Thus, cardiac and (cerebro-) vascular pathology affecting CBF might influence cellular functions in the brain before structures are altered. Exploring the contribution of cardiac and (cerebro-)vascular pathology to brain alterations is important, because it could identify treatment targets for patients with cognitive impairment due to this type of pathology in the foreseeable future. Medication that improves hemodynamics, such as antihypertensive medication, is available at this moment. However, trials assessing the efficacy of such therapies in VCI are currently lacking. This is due to an incomplete understanding of the mechanisms involved and to the lack of research that identifies patients who will benefit most [1, 16, 17]. Since health care and research are usually





Cerebrovasc Dis Extra 2017;7:140–152

Hooghiemstra et al.: Design of the Heart-Brain Study

Textbox 1

The Heart-Brain Connection consortium

The Heart-Brain Connection consortium consists, in addition to the Heart-Brain Study, of preclinical, experimental, and clinical trial research. Here we briefly describe these studies and how they are connected to the clinical study.

- 1 Within the data of the Rotterdam Study [58], a large prospective cohort study in the city of Rotterdam, the Netherlands, we investigate the heart-brain link from a population-based perspective. If we can demonstrate putative causal associations, this extends the importance of cardiac function and cerebral hemodynamics for brain disease to an asymptomatic population.
- 2 In a preclinical program, we focus on the senescence accelerated mouse (SAMP8) model to test the link between cardiovascular function, cerebral blood flow, and cognition in combination with potential genetic aggravating factors [59–62]. This mouse model shows cardiac hypertrophy as well as heart failure. This study advances our insight into the importance of hemodynamics for brain function, as well as the interplay between systemic hypoperfusion and disturbed local vessel function.
- 3 We perform an additional observational imaging study performed on a 7-tesla MRI scanner. With this ultra-high-field MRI we evaluate novel techniques assessing intracranial small vessel pulsatility and microvascular functioning to explore their value as novel etiologic, diagnostic, and prognostic biomarkers for VCI.
- 4 We conduct a randomized controlled trial aimed at improving cerebral perfusion (as measured with MRI-based arterial spin labeling) in elderly patients with VCI through aerobic exercise, while taking into account a potential modulatory effect of cardiac output [63]. This proof-of-principle RCT may show that improvement of the cerebral perfusion can lead to improved cognitive functioning in patients with VCI.

MRI, magnetic resonance imaging; RCT, randomized, controlled trial; SAMP8, senescence accelerated mouse; VCI, vascular cognitive impairment.

organized in a monodisciplinary way, cardiovascular status tends to be neglected in patients presenting with cognitive impairment in memory clinics and, vice versa, cognitive disorders are often neglected in patients presenting with cardiovascular disease in cardiology or vascular medicine departments. Moreover, guidelines for diagnostic protocols that provide a combined comprehensive assessment of the cardiovascular and cerebral structure and function are lacking. The Heart-Brain Study is part of a larger Heart-Brain Connection consortium (www. heart-brain.nl) [18], covering preclinical, experimental, and clinical trial research (Textbox 1). The Heart-Brain Study hypothesizes that the hemodynamic status of the heart and the brain is an important, but underestimated cause of VCI. We aim to assess the association between (cardio-)vascular and hemodynamic factors in the heart and the brain in relation to cognitive function. Our objectives are to assess (1) the association between cardiovascular parameters and cognitive function, (2) the association between cardiovascular parameters and brain structure and perfusion, and (3) the association between brain structure and perfusion and cognitive function. We study these objectives in patients with HF, COD, and VCI, both crosssectionally as well as longitudinally. By studying the complete heart-brain axis in patient groups that present themselves with complaints attributed to specific parts of the heart-brain axis, we have the opportunity to assess a combination of clinical and subclinical manifestations of disorders of the heart, vascular system, and brain, with hemodynamic status as a possible binding factor. Here, we describe the design and study protocol of this multidisciplinary study, in which cardiologists, epidemiologists, neurologists, neuropsychologists, radiologists, image processing experts, and MR physicists work together to study the hemodynamic status of the heart and the brain as an important, but underestimated determinant of VCI.





Cerebrovasc Dis Extra 2017;7:140–152	
DOI: 10.1159/000480738	$\ensuremath{\mathbb{C}}$ 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Tahlo	1 (Overview	of data	collection	ner	assessment moment
lable	- . '		uata u	conection	per	assessment moment

	Baseline	Follow-up 1 year ^a	Follow-up 2 years
Informed consent	Х		
Clinical assessment	Х		Х
Neuropsychological assessment	Х	Х	Х
Brain MRI	Х		Х
Cardiac MRI	Х		
Blood	Х		Х
Cerebrospinal fluid ^b	Х		
Echocardiography ^c	Х		Х
24-hour blood pressure ^d	Х		
Cause of death for deceased patients		Х	Х

^a Only for patients with vascular cognitive impairment; ^b Performed in the routine clinical setting in VU University Medical Center; ^c Available in Maastricht University Medical Center and VU University Medical Center; ^d Available in Maastricht University Medical Center, University Medical Center Utrecht, and VU University Medical Center.

Methods

Study Design

The Heart-Brain Study is a prospective study with a follow-up measurement after 2 years. Five Dutch university medical centers collaborate: Erasmus Medical Center (ErasmusMC) in Rotterdam, Leiden University Medical Center (LUMC) in Leiden, Maastricht University Medical Center (MUMC) in Maastricht, University Medical Center Utrecht (UMCU) in Utrecht, and VU University Medical Center (VUmc) in Amsterdam. Participants have been enrolled from September 2014 onwards. For an overview of the data collection see Table 1.

Participants

Patients with VCI, COD, and HF are recruited from cardiology, memory, and neurology outpatient clinics from four sites: LUMC, MUMC, UMCU, and VUmc. In each patient group, 175 patients are recruited, yielding a total sample size of 525 patients. In addition, 120 controls undergo the same study procedures as patients. Eligible participants are selected according to the inclusion and exclusion criteria (Table 2). Written informed consent is obtained prior to participation in the study.

Baseline Assessment

Clinical Data and Assessment

The following measures are collected and saved as a core clinical dataset:

- Sociodemographic factors, including age, sex, educational level, and social situation.
- Vascular risk factors including hypertension, diabetes, hyperlipidemia, smoking, overweight, and extensive alcohol use.
- Medical, neurologic, cardiovascular, and family history.
- Current medication.
- Physical examination with particular attention to clinical signs of volume overload (e.g., pitting edema, rales) and heart murmur.
- Duplicate blood pressure measurement on one occasion (sitting, lying, and standing).
- Resting 12-lead electrocardiography.
- Anthropometry, physical performance, and physical activity [19, 20].





Cerebrovasc Dis Extra 2017;7:14	0–152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Table 2. Inclusion and exclusion criteria of patients with VCI. COD, and HF and controls for the Heart-Brain Study

General selection criteria for all patient groups Inclusion criteria

Age 50 years or older

- Able to undergo cognitive testing
- Independence in daily life

Exclusion criteria

Contraindication for MRI or unable to undergo MRI protocol due to physical condition _

- Life-threatening disease with life expectancy less than 3 years other than VCI, COD, or HF
- A clinical diagnosis of dementia is not a contraindication for participation in this study; however, clinical evidence of a neurodegenerative disease other than VCI or AD (such as frontotemporal dementia, Lewy body disease, or hypokinetic rigid syndrome) is an exclusion criterion
- Another neurologic or psychiatric diagnosis that affects cognitive performance or testing, such as severe traumatic brain injury or substance abuse
- Participation in ongoing trials for therapeutic interventions including randomized controlled trials and clinical trials of investigational medicinal products
- Plan to move out of the region within the next 3 years
- Atrial fibrillation at the moment of inclusion (of note, [paroxysmal] atrial fibrillation in the history is not an exclusion criterion). PVCs exceeding 10% of the total number of heartbeats, e.g., a heart rate of 60/min and >6 PVCs

Additional selection criteria for patients with VCI Inclusion criteria

- Cognitive complaints
- Clinical Dementia Rating ≤1 and Mini-Mental State Examination ≥20
- Furthermore, at least one of the following criteria should be present:
- On brain MRI moderate to severe white matter lesion (Fazekas >1) and/or (lacunar) infarct(s) and/or intracerebral (micro-)hemorrhage(s)
- On brain MRI mild white matter lesions (Fazekas = 1) and at least two of the following vascular risk factors: hypertension, hypercholesterolemia, diabetes mellitus, obesity, smoking, or clinically manifest vascular disease (>6 months ago). Clinically manifest vascular disease comprises peripheral arterial disease, myocardial infarction, percutaneous coronary intervention/coronary artery bypass graft, and/or stroke

Exclusion criteria

n/a

Additional selection criteria for patients with COD Inclusion criterion

- Significant stenosis (>80%) or occlusion of the internal carotid artery as visible on MR angiography Exclusion criterion
 - Plan for carotid surgery

Additional selection criteria for patients with HF

Inclusion criteria

This study includes HF patients irrespective of left ventricular ejection fraction and coronary artery disease HF according to European Cardiology Society guidelines:

- Symptoms typical of HF (breathlessness at rest or on exercise, fatigue, tiredness, ankle swelling)
- Signs typical of HF (tachycardia, tachypnea, pulmonary rales, pleural effusion, raised jugular venous pressure, peripheral edema, hepatomegaly)
- Objective evidence of a structural or functional abnormality of the heart at rest on routine echocardiography
- Stable clinical situation for at least 6 months

Exclusion criteria

n/a

Additional selection criteria for controls

Inclusion criteria

n/a

Exclusion criterion

A diagnosis of VCI, COD, and HF, i.e., a control may participate when one or two diagnoses are present

AD, Alzheimer disease; COD, carotid occlusive disease; HF, heart failure; MRI, magnetic resonance imaging; PVC, premature ventricular contraction; VCI, vascular cognitive impairment.



Cerebrovasc Dis Extra 2017;7:140	0–152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Test/questionnaire	Domain
Cognitive functioning	
Mini-Mental State Examination (MMSE) [44]	Global cognition
15-Word Auditory Verbal Learning Test [45]	Episodic memory
Total recall	
Delayed recall	
Recognition	
Visual Association Test (VAT), short version [46]	Implicit associative visual learning
Digit Span, extended version [47]	
Forward	Attention
Backward	Working memory
Trail Making Test (TMT) [48]	
Part A	Information processing speed, attention
Part B	Response inhibition, executive functioning
Stroop Color-Word Test (SCWT) [49]	
Card I and II	Information processing speed, attention
Card III	Concept shifting, executive functioning
Fluency, 60 s (animals) [50, 51]	Verbal fluency, semantic memory
Letter-Digit Substitution Test (LDST) [52]	Information processing speed, attention
General functioning and neuropsychiatry	
Clinical Dementia Rating Scale (CDR) [53]	Severity of cognitive impairment
Amsterdam IADL Questionnaire [54]	Functional status
Disability Assessment of Dementia (DAD) [55]	Functional status
Geriatric Depression Scale-15 (GDS-15) [56]	Depressive symptoms
Starkstein Apathy Scale [57]	Symptoms of apathy

Table 3. Neuropsychological assessment and measures of neuropsychiatry and general functioning

CDR, Clinical Dementia Rating scale; DAD, Disability Assessment Dementia; GDS-15, Geriatric Depression Scale-15; IADL, Instrumental Activity of Daily Living; LDST, Letter-Digit Substitution Test; MMSE, Mini-Mental State Examination; TMT, Trail Making Test; SCWT, Stroop Color-Word Test; VAT, Visual Association Test.

Cognitive Functioning

All participants undergo an extensive and standardized neuropsychological assessment, based on the Dutch Parelsnoer Initiative [21]. This test battery covers global cognitive functioning and four major cognitive domains including memory, language, attention-psychomotor speed, and executive functioning (Table 3). All test scores are standardized into *z*-scores and subsequently combined into cognitive domains. We rate a cognitive domain as impaired when the *z*-score is below -1.5. Patients are classified as cognitively normal when no domains are impaired, with minor cognitive impairment with one domain impaired, and with major cognitive impairment with more than one domain impaired.

In addition to cognitive functioning, we assess general functioning, activities of daily living, depressive symptoms, and apathy (Table 3).

MRI

KARGER

MRIs are acquired on Philips Ingenia 3T scanners at LUMC and UMCU, a Philips Achieva 3T scanner at MUMC, and a Philips Gemini 3T PET-MR scanner at VUmc (Philips, Best, The Netherlands). The MRI protocol consists of a cardiac, vascular, and brain protocol (Table 4). The brain protocol includes T1-weighted, fluid-attenuated inversion recovery (FLAIR) images and susceptibility-weighted imaging (SWI). Cerebral perfusion is measured with arterial spin labeling [22] and phase-contrast flow measurements. The sequences measuring perfusion



Cerebrovasc Dis Extra 2017;7:1	40–152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Table 4. MRI protocol

Organ	Pathophysiological phenomenon	Parameter	MR technique	Duration, min:s	Resolution, mm ³	Relevant contrast parameters
Brain	Structural status	Atrophy (brain volumes)	1. T1-weighted	6:47	1×1×1	MP-RAGE; TR 8.2 ms; TE 4.5 ms; shot interval 3,000 ms; flip angle 8°; inversion delay 990 ms
		WMH + infarcts	2. FLAIR	4:43	1.11×1.11×1.11	TR 4,800 ms; TE 313 ms; TI 1,650 ms; TSE factor 182
		Microbleeds	3. SWI	2:30	0.8×0.8×1.6	3D gradient echo; TR 45 ms; TE 31 ms; flip angle 13°; EPI factor 3
	Perfusion at rest	Whole brain perfusion at rest	4. ASL	6:05	3×3×7	pCASL; label duration 1,800 ms; postlabeling delay 1,800 ms; background suppression; multislice 2D; single shot EPI readout
	Cerebral blood flow	Total cerebral blood flow	5. Phase-contrast flow measurement	0:43	1.17×1.17×5	TR 12 ms; TE 8.2 ms; flip angle 10°; Venc 200 cm/s; untriggered; 10 averages
Aorta	Aorta stiffness	Pulse wave velocity	6. Aorta QFlow	1:47	2.5×2.5×8	TR 4.7 ms; TE 2.8 ms; flip angle 10°; Venc 150 cm/s; number of heart phases dependent on heart rate; temporal resolution 5 ms
Heart	Functional status	Systolic function	7. Short-axis multi-slice cine SSFP	3:00	1.5×1.6×8.0	TR 3.1 ms; TE 1.55; flip angle 45°; 40 heart phases; 67 phase percentage; breath-hold; number of slices dependent on size of LV (range 12–16 slices)
	Structural status	Diastolic function Cardiac output LV mass LV volume	8. Phase contrast mitral inflow Comes with 7 Comes with 7 Comes with 7	2:00	2.5×2.5×8.0	TR 4.4 ms; TE 2.8 ms; flip angle 10°; 40 heart phases; Venc 150 cm/s

ASL, arterial spin labeling; EPI factor, echo-planar imaging factor; FLAIR, fluid attenuation inversion recovery; LV, left ventricular; MP-RAGE, magnetizationprepared rapid acquisition gradient echo; pCASL, pseudo-continuous arterial spin labeling; Qflow, quantative flow; SSFP, steady-state free precession; SWI, susceptibility-weighted imaging; TE, echo time; TI, inversion time; TR, repetition time; TSE factor, turbo spin-echo factor; Venc, velocity encoding; WMH, white matter hyperintensities.

are performed in the same scan session as the structural sequences. The heart protocol includes short-axis multislice cine steady-state free precision (SSFP), aorta QFlow images, and phase-contrast mitral flow measurements. Scans are screened by local radiologists for clinically relevant incidental findings.

Blood and Cerebrospinal Fluid Markers

We investigate systemic and organ-specific blood biomarkers that relate to functional or structural abnormalities in components of the heart-brain axis. For systemic biomarkers, we assess biomarkers related to processes involved in HF, atherosclerosis, and VCI. We focus on abnormalities in lipid metabolism, insulin resistance/dysglycemia, inflammation, and anemia. For markers reflecting pathogenic processes in organ-specific components of the heart-blood vessels-brain axis, we assess markers of HF and cardiac fibrosis, and remodeling of blood vessel pathology and of Alzheimer-type pathology.

Cerebrospinal fluid is collected for patients with VCI in the routine clinical setting in VUmc for determination of amyloid-beta 1–42, total tau, and hyperphosphorylated tau-18 [23]. Participants are asked to give separate informed consent for DNA storage for future genetic analyses.

Echocardiography

We use standard clinical Doppler-echocardiographic equipment to measure the complete standard clinical array including structures as well as systolic and diastolic function of both ventricles, atrial dimensions, and valve function, as recommended by the European and



D' E + 00177140 150



Cerebrovasc Dis Extra 2017,7.14	0-152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Hooghiemstra et al.: Design of the Heart-Brain Study

American Societies of Echocardiography [24, 25]. Transthoracic echocardiography is performed in standard parasternal, apical, and subcostal views. Echocardiography is performed in MUMC and VUmc.

24-Hour Ambulatory Blood Pressure Monitoring

24-hour ambulatory blood pressure monitoring is performed, using validated blood pressure monitors from Microlife (Microlife Corporation Europe, Widnau, Switzerland). 24-hour blood pressure measurements have been shown to be of better prognostic value for cardiovascular events and more reproducible than conventional office BP measurements [26]. 24-hour ambulatory blood pressure is performed in MUMC, UMCU, and VUmc.

Follow-Up Assessment

Two years after baseline assessment, all participants are invited for a second visit which includes neuropsychological testing, brain MRI, and collection of blood samples. Echocardiography is included when performed at baseline. In addition, clinical data on disease incidence and admission to hospital or nursing home between the first and last visit are collected by history taking. For participants with cognitive problems or when the history of the participant is considered less reliable, additional history is obtained from next of kin and/or the general practitioner. For deceased participants, the cause of death is obtained from Central Agency for Statistics Netherlands (CBS) and general practitioners. Patients with VCI additionally undergo follow-up of neuropsychological testing after 1 year. We evaluate cognitive decline based on the difference in the cognitive domain z-scores (for more information, see section Cognitive Functioning above).

Data Collection, Processing, and Storage

We use centralized state-of-the-art storage platforms for clinical (OpenClinica, LLC, Waltham, MA, USA) and imaging data (Extensible Neuroimaging Archive Toolkit [XNAT]). Imaging data are processed centrally with automated standardized pipelines. For an elaborate description, see online supplementary material (for all online suppl. material, see www. karger.com/doi/10.1159/000480738).

Sample Size Considerations

With a sample size of 175 patients in each patient cohort we can detect associations in which the determinant explains 4% or more of the variance in the dependent variable (i.e., the equivalent of a correlation coefficient of 0.2 or more with alpha 0.05, power 90%), taking 10–20 relevant covariates into account. Assuming that the dropout rate will not exceed 25% over 2 years, we have 80% power to detect associations of the same strength at follow-up.

Statistical Analysis

Cross-Sectional Relations

Regression analyses are used to investigate the independent associations between measures of cerebral perfusion and blood flow, structural brain abnormalities (brain atrophy, white matter hyperintensities, infarcts, and microbleeds), and cognitive performance. Also, regression analyses are used to investigate the relationship between cardiovascular parameters (cardiac output, systolic and diastolic function of the ventricle, blood pressure, pulse wave velocity, aortic and carotid stiffness) and cerebral perfusion and blood flow (arterial spin labeling and phase-contrast flow measurements) at baseline.

KARGER



DOI: 10.1159/000480738 © 2017 The Author(s). Published by S. Karger AG www.karger.com/cee	i, Base

Prospective Relations

To investigate prospective relations of baseline cardiovascular and cerebral perfusion and flow measures with change in brain MRI abnormalities and cognitive functioning, we use regression models with brain volume and change in cognitive performance at follow-up as the dependent variables and cardiovascular parameters and brain perfusion and blood flow at baseline as the independent variables.

Since the main analyses have an etiologic focus, appropriate adjustment for confounding factors is performed. All abovementioned associations are examined in each patient cohort (VCI, COD, or HF) separately, comparing groups with controls. Finally, we pool all data and perform linear regression analyses, taking into account potential effect modifications by cohort [27].

Ethical Considerations

The Medical Ethics Review Committee of the LUMC performed central approval of the Heart-Brain Study (number P.14.002). Subsequently, local boards of the UMCs approved the local performance of the study. The Heart-Brain Study is performed in accordance with the declaration of Helsinki (version 2013) and the Medical Research Involving Human Subjects Act (WMO).

Current Status and Time Line

The first participant was included in September 2014, the inclusion period finalizes in 2017, and the last follow-up measurement will be in 2018. The first baseline results are expected in 2017, the longitudinal results in 2019.

Results and Conclusion

The Heart-Brain Study hypothesizes that the hemodynamic status of the heart and the brain are important, but underestimated determinants of VCI. Previous studies have investigated components of the heart-brain axis in (prospective cohorts of) healthy people [28-33], patients with HF [12, 13, 34–39], and COD [40–43]. These studies have found circumstantial evidence that cardiac and cardiovascular pathology affecting CBF and perfusion in the brain may influence brain function before structures are irreversibly damaged. It is currently unknown how often hemodynamic changes based on cardiovascular pathology occur in patients with cognitive impairment. Various cardiovascular factors, such as cardiac output, blood pressure, pulse wave velocity, and aortic and carotid stiffness, may influence CBF. On the other hand, in VCI a lower CBF could also be related to a decreased need of blood by an already affected brain. Little is known about how these factors, separate or in concert, influence cognitive performance. The Heart-Brain Study is unique because of the integrated approach that we use to investigate relationships between (cardio-)vascular factors, the hemodynamic status of the heart and the brain, and cognitive impairment, in three patient groups that represent components of the heart-brain axis. While zooming in on one component of the heart-brain axis we assess the other components and how they are interconnected. This way, we assess both clinical and subclinical manifestations of disorders of the heart, vascular system, and brain, with hemodynamic status as a possible binding factor along the heartbrain axis. This integrated approach may show light on the mechanisms involved in these relationships. To study the relationships as clearly as possible we chose to exclude patients with current atrial fibrillation at the time of inclusion, since atrial fibrillation may lead to unpredictable hemodynamic changes. This exclusion might lead to limited generalizability of this study. However, the patient groups mainly function as a model of specific parts of the





Cerebrovasc Dis Extra 2017;7:1	40–152
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

heart-brain axis, i.e., hemodynamic components possibly leading to chronic cerebral hypoperfusion.

We perform extensive phenotyping using a comprehensive and standardized MRI protocol that has been developed to measure structure and function of both the heart and the brain. Alongside, a platform for data storage and image processing is developed in which both automatic and manual quality assessment procedures are implemented. Quantification of imaging biomarkers of the heart, brain, and cerebropetal arteries is performed with existing and newly developed automated software. With this study, we provide a foundation for an interdisciplinary collaborative network for the study of the heart-brain axis that will lead to a true multidisciplinary and consensus-based approach of clinical management of cognitive impairment in patients with HF, COD, and VCI. The close collaboration between departments of cardiology and neurology opens possibilities for future heart-brain clinics, through which implementation of newly developed diagnostic tools and treatment options can be optimized. With this approach we meet the clinical and research need for centers of excellence with transdisciplinary programs within and between centers [1, 16, 17].

In addition to the Heart-Brain Study, in the Heart-Brain Connection consortium [18] we perform preclinical, experimental, and clinical trial studies that further increase the understanding of the mechanisms underlying the relationship between hemodynamic status and cognitive functioning (Textbox 1).

In conclusion, in the Heart-Brain Study we test the hypothesis that the hemodynamic status of the heart and the brain is an important, but underestimated cause of VCI offering promising opportunity for treatment. Moreover, we develop a novel, clinically feasible diagnostic protocol including a comprehensive MRI protocol that assesses the heart, the vascular system, and the brain. This protocol can be used for identifying patients suitable for future trials as well as monitoring treatment effects. Finally, we provide a foundation for an inter-disciplinary collaboration for the study of VCI that will lead to a true multidisciplinary and consensus-based approach of the clinical management of VCI.

Acknowledgements

Cerebrovascular Diseases

We acknowledge the contribution of Heart-Brain Study researchers and employees. For an overview of Heart-Brain Connection consortium members, see online supplementary material.

We acknowledge the support of the Netherlands CardioVascular Research Initiative: the Dutch Heart Foundation (CVON 2012-06 Heart Brain Connection), Dutch Federation of University Medical Centers, the Netherlands Organization for Health Research and Development, and the Royal Netherlands Academy of Sciences.

Disclosure Statement

KARGER

W.J. Niessen is cofounder, part-time Chief Scientific Officer, and stock holder of Quantib BV. Other authors declare that there are no competing interests. None of the authors have direct or indirect relationships with the Netherlands CardioVascular Research Initiative.





DOI: 10.1159/000480738 © 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

Hooghiemstra et al.: Design of the Heart-Brain Study

References

- 1 Gorelick PB, Scuteri A, Black SE, Decarli C, Greenberg SM, Iadecola C, et al: Vascular contributions to cognitive impairment and dementia: a statement for healthcare professionals from the American Heart Association/ American Stroke Association. Stroke 2011;42:2672–2713.
- 2 Ganguli M: Epidemiology of Dementia; in Principles and Practice of Geriatric Psychiatry. Hoboken, Wiley, 2010, pp 207–212.
- 3 Lobo A, Launer LJ, Fratiglioni L, Andersen K, Di Carlo A, Breteler MM, et al: Prevalence of dementia and major subtypes in Europe: a collaborative study of population-based cohorts. Neurology 2000;54:4–9.
- 4 Ferri CP, Prince M, Brayne C, Brodaty H, Fratiglioni L, Ganguli M, et al: Global prevalence of dementia: a Delphi consensus study. Lancet 2005;366:2112–2117.
- 5 Hachinski V, Iadecola C, Petersen RC, Breteler MM, Nyenhuis DL, Black SE, et al: National Institute of Neurological Disorders and Stroke-Canadian Stroke Network vascular cognitive impairment harmonization standards. Stroke 2006;37:2220–2241.
- 6 Sachdev P, Kalaria R, O'Brien J, Skoog I, Alladi S, Black SE, et al: Diagnostic criteria for vascular cognitive disorders: a VASCOG statement. Alzheimer Dis Assoc Disord 2014;28:206–218.
- 7 O'Brien JT, Erkinjuntti T, Reisberg B, Roman G, Sawada T, Pantoni L, et al: Vascular cognitive impairment. Lancet Neurol 2003;2:89–98.
- 8 Skrobot OA, O'Brien JO, Black S, Chen C, DeCarli C, Erkinjuntti T, et al: The Vascular Impairment of Cognition Classification Consensus Study. Alzheimers Dement 2017;13:624–633.
- 9 Marshall RS, Lazar RM: Pumps, aqueducts, and drought management: vascular physiology in vascular cognitive impairment. Stroke 2011;42:221–226.
- 10 Meissner A: Hypertension and the brain: a risk factor for more than heart disease. Cerebrovasc Dis 2016;42: 255–262.
- 11 Okonkwo OC, Cohen RA, Gunstad J, Tremont G, Alosco ML, Poppas A: Longitudinal trajectories of cognitive decline among older adults with cardiovascular disease. Cerebrovasc Dis 2010;30:362–373.
- 12 Bornstein RA, Starling RC, Myerowitz PD, Haas GJ: Neuropsychological function in patients with end-stage heart failure before and after cardiac transplantation. Acta Neurol Scand 1995;91:260–265.
- 13 Zuccala G, Onder G, Marzetti E, Monaco MR, Cesari M, Cocchi A, et al: Use of angiotensin-converting enzyme inhibitors and variations in cognitive performance among patients with heart failure. Eur Heart J 2005;26: 226–233.
- 14 Roman DD, Kubo SH, Ormaza S, Francis GS, Bank AJ, Shumway SJ: Memory improvement following cardiac transplantation. J Clin Exp Neuropsychol 1997;19:692–697.
- 15 Sasoh M, Ogasawara K, Kuroda K, Okuguchi T, Terasaki K, Yamadate K, et al: Effects of EC-IC bypass surgery on cognitive impairment in patients with hemodynamic cerebral ischemia. Surg Neurol 2003;59:453–455.
- 16 Snyder HM, Corriveau RA, Craft S, Faber JE, Knopman D, Lamb BT, et al: Vascular contributions to cognitive impairment and dementia including Alzheimer's disease. Alzheimers Dement 2015;11:710–717.
- 17 Daviglus M, Bell C, Berrettini W, Bowen P, Connolly E, Cox N, et al: NIH state-of-the-science conference statement: preventing Alzheimer's disease and cognitive decline. NIH Consens State Sci Statements 2010;27: 1–30.
- 18 van Buchem MA, Biessels GJ, Brunner la Rocca HP, de Craen AJM, van der Flier WM, Ikram MA, et al: The heartbrain connection: a multidisciplinary approach targeting a missing link in the pathophysiology of vascular cognitive impairment. J Alzheimers Dis 2014;42(suppl 4):S443–S451.
- 19 Guralnik JM, Simonsick EM, Ferrucci L, Glynn RJ, Berkman LF, Blazer DG, et al: A short physical performance battery assessing lower extremity function: association with self-reported disability and prediction of mortality and nursing home admission. J Gerontol 1994;49:85–94.
- 20 Pols MA, Peeters PH, Ocké MC, Slimani N, Bueno-de-Mesquita HB, Collette HJ: Estimation of reproducibility and relative validity of the questions included in the EPIC Physical Activity Questionnaire. Int J Epidemiol 1997;26:181–189.
- 21 Aalten P, Ramakers IHGB, Biessels GJ, de Deyn PP, Koek HL, OldeRikkert MGM, et al: The Dutch Parelsnoer Institute-Neurodegenerative diseases; methods, design and baseline results. BMC Neurol 2014;14:254.
- 22 Alsop DC, Detre JA, Golay X, Günther M, Hendrikse J, Hernandez-Garcia L, et al: Recommended implementation of arterial spin-labeled perfusion MRI for clinical applications: a consensus of the ISMRM perfusion study group and the European consortium for ASL in dementia. Magn Reson Med 2015;73:102–116.
- 23 Duits FH, Teunissen CE, Bouwman FH, Visser PJ, Mattsson N, Zetterberg H, et al: The cerebrospinal fluid "Alzheimer profile": Easily said, but what does it mean? Alzheimers Dement 2014;10:713–723.
- 24 Quiñones MA, Otto CM, Stoddard M, Waggoner A, Zoghbi WA: Recommendations for quantification of Doppler echocardiography: a report from the Doppler Quantification Task Force of the Nomenclature and Standards Committee of the American Society of Echocardiography. J Am Soc Echocardiogr 2002;15:167–184.
- 25 Lang RM, Badano LP, Mor-Avi V, Afilalo J, Armstrong A, Ernande L, et al: Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. J Am Soc Echocardiogr 2015;28:1–39.
- 26 Parati G, Stergiou GS, Asmar R, Bilo G, de Leeuw P, Imai Y, et al: European Society of Hypertension guidelines for blood pressure monitoring at home: a summary report of the Second International Consensus Conference on Home Blood Pressure Monitoring. J Hypertens 2008;26:1505–1526.





- 27 Rothman K, Greenland S, Lash T: Modern Epidemiology, ed 3 (mid-cycle revision). Philadelphia, Wolters Kluwer/Lippincott WIliams & Wilkins, 2016.
- 28 Jefferson AL, Himali JJ, Beiser AS, Au R, Massaro JM, Seshadri S, et al: Cardiac index is associated with brain aging: the Framingham Heart Study. Circulation 2010;122:690–697.
- 29 Mitchell GF, van Buchem MA, Sigurdsson S, Gotal JD, Jonsdottir MK, Kjartansson O, et al: Arterial stiffness, pressure and flow pulsatility and brain structure and function: the Age, Gene/Environment Susceptibility-Reykjavik study. Brain 2011;134:3398–3407.
- 30 Muller M, Grobbee DE, Aleman A, Bots M, van der Schouw YT: Cardiovascular disease and cognitive performance in middle-aged and elderly men. Atherosclerosis 2007;190:143–149.
- 31 Scuteri A, Brancati AM, Gianni W, Assisi A, Volpe M: Arterial stiffness is an independent risk factor for cognitive impairment in the elderly: a pilot study. J Hypertens 2005;23:1211–1216.
- 32 Elias MF, Robbins MA, Budge MM, Abhayaratna WP, Dore GA, Elias PK: Arterial pulse wave velocity and cognition with advancing age. Hypertension 2009;53:668–673.
- 33 Waldstein SR, Rice SC, Thayer JF, Najjar SS, Scuteri A, Zonderman AB: Pulse pressure and pulse wave velocity are related to cognitive decline in the Baltimore Longitudinal Study of Aging. Hypertension 2008;51:99–104.
- 34 Paulson OB, Jarden JO, Godtfredsen J, Vorstrup S: Cerebral blood flow in patients with congestive heart failure treated with captopril. Am J Med 1984;76:91–95.
- 35 Vogels RLC, Oosterman JM, van Harten B, Scheltens P, van der Flier WM, Schroeder-Tanka JM, et al: Profile of cognitive impairment in chronic heart failure. J Am Geriatr Soc 2007;55:1764–1770.
- 36 Vogels RLC, van der Flier WM, van Harten B, Gouw AA, Scheltens P, Schroeder-Tanka JM, et al: Brain magnetic resonance imaging abnormalities in patients with heart failure. Eur J Heart Fail 2007;9:1003–1009.
- 37 Vogels RLC, Oosterman JM, van Harten B, Gouw AA, Schroeder-Tanka JM, Scheltens P, et al: Neuroimaging and correlates of cognitive function among patients with heart failure. Dement Geriatr Cogn Disord 2007;24:418– 423.
- 38 Vogels RLC, Oosterman JM, Laman DM, Gouw AA, Schroeder-Tanka JM, Scheltens P, et al: Transcranial Doppler blood flow assessment in patients with mild heart failure: correlates with neuroimaging and cognitive performance. Congest Heart Fail 2008;14:61–65.
- 39 Choi B-R, Kim JS, Yang YJ, Park K-M, Lee CW, Kim Y-H, et al: Factors associated with decreased cerebral blood flow in congestive heart failure secondary to idiopathic dilated cardiomyopathy. Am J Cardiol 2006;97:1365– 1369.
- 40 Bakker FC, Klijn CJM, Jennekens-Schinkel A, van der Tweel I, Tulleken CAF, Kappelle LJ: Cognitive impairment in patients with carotid artery occlusion and ipsilateral transient ischemic attacks. J Neurol 2003;250:1340– 1347.
- 41 Bakker FC, Klijn CJM, Jennekens-Schinkel A, van der Tweel I, van der Grond J, van Huffelen AC, et al: Cognitive impairment is related to cerebral lactate in patients with carotid artery occlusion and ipsilateral transient ischemic attacks. Stroke 2003;34:1419–1424.
- 42 Bakker FC, Klijn CJM, van der Grond J, Kappelle LJ, Jennekens-Schinkel A: Cognition and quality of life in patients with carotid artery occlusion: a follow-up study. Neurology 2004;62:2230–2235.
- 43 Frazier DT, Seider T, Bettcher BM, Mack WJ, Jastrzab L, Chao L, et al: The role of carotid intima-media thickness in predicting longitudinal cognitive function in an older adult cohort. Cerebrovasc Dis 2014;38:441–447.
- 44 Folstein MF, Folstein SE, Mchugh PR: "Mini-mental state." A practical method for grading the cognitive state of patients for the clinician. J Psychiatr Res 1975;12:189–198.
- 45 Van der Elst W, van Boxtel MPJ, van Breukelen GJP, Jolles J: Rey's verbal learning test: normative data for 1855 healthy participants aged 24–81 years and the influence of age, sex, education, and mode of presentation. J Int Neuropsychol Soc 2005;11:290–302.
- 46 Lindeboom J, Schmand B, Tulner L, Walstra G, Jonker C: Visual association test to detect early dementia of the Alzheimer type. J Neurol Neurosurg Psychiatry 2002;73:126–133.
- 47 Lindeboom J, Matto D: Digit series and Knox cubes as concentration tests for elderly subjects. Tijdschr Gerontol Geriatr 1994;25:63–68.
- 48 Reitan RM: The relation of the trail making test to organic brain damage. J Consult Psychol 1955;19:393–394.
- 49 Van der Elst W, Van Boxtel MPJ, Van Breukelen GJP, Jolles J: The Stroop color-word test: influence of age, sex, and education; and normative data for a large sample across the adult age range. Assessment 2006;13:62–79.
- 50 Van der Elst W, Van Boxtel MPJ, Van Breukelen GJP, Jolles J: Normative data for the Animal, Profession and Letter M Naming verbal fluency tests for Dutch speaking participants and the effects of age, education, and sex. J Int Neuropsychol Soc 2006;12:80–89.
- 51 Luteijn F, van der Ploeg FAE: Handleiding Groninger Intelligentie Test (Manual Groningen Intelligence Test). Lisse, Swets & Zeitlinger, 1983.
- 52 van der Elst W, van Boxtel MPJ, van Breukelen GJP, Jolles J: The Letter Digit Substitution Test: normative data for 1,858 healthy participants aged 24–81 from the Maastricht Aging Study (MAAS): influence of age, education, and sex. J Clin Exp Neuropsychol 2006;28:998–1009.
- 53 Morris JC: Clinical dementia rating: a reliable and valid diagnostic and staging measure for dementia of the Alzheimer type. Int Psychogeriatr 1997;9:173–178.
- 54 Sikkes S, de Lange-de Klerk E, Pijnenburg Y, Gillissen F, Romkes R, Knol DL, et al: A new informant-based questionnaire for instrumental activities of daily living in dementia. Alzheimers Dement 2012;8:536–543.





Cerebrovasc Dis Extra 2017;7:140–152	
DOI: 10.1159/000480738	© 2017 The Author(s). Published by S. Karger AG, Basel www.karger.com/cee

- 55 Gelinas I, Gauthier L, McIntyre M, Gauthier S: Development of a functional measure for persons with Alzheimer's disease: the disability assessment for dementia. Am J Occup Ther 1999;53:471–481.
- 56 Yesavage JA, Brink TL, Rose TL, Lum O, Huang V, Adey M, et al: Development and validation of a geriatric depression screening scale: a preliminary report. J Psychiatr Res 1983;17:37–49.
- 57 Starkstein SE, Jorge R, Mizrahi R, Robinson RG: A prospective longitudinal study of apathy in Alzheimer's disease. J Neurol Neurosurg Psychiatry 2006;77:8–11.
- 58 Ikram MA, van der Lugt A, Niessen WJ, Koudstaal PJ, Krestin GP, Hofman A, et al: The Rotterdam Scan Study: design update 2016 and main findings. Eur J Epidemiol 2015;30:1299–1315.
- 59 Poulet R, Gentile MT, Vecchione C, Distaso M, Aretini A, Fratta L, et al: Acute hypertension induces oxidative stress in brain tissues. J Cereb Blood Flow Metab 2006;26:253–262.
- 60 Carnevale D, Mascio G, Ajmone-Cat MA, D'Andrea I, Cifelli G, Madonna M, et al: Role of neuroinflammation in hypertension-induced brain amyloid pathology. Neurobiol Aging 2012;33:19–29.
- 61 Gentile MT, Poulet R, Pardo AD, Cifelli G, Maffei A, Vecchione C, et al: Beta-amyloid deposition in brain is enhanced in mouse models of arterial hypertension. Neurobiol Aging 2009;30:222–228.
- 62 Carnevale D, Mascio G, D'Andrea I, Fardella V, Bell RD, Branchi I, et al: Hypertension induces brain betaamyloid accumulation, cognitive impairment, and memory deterioration through activation of receptor for advanced glycation end products in brain vasculature. Hypertension 2012;60:188–197.
- 63 Leeuwis AE, Hooghiemstra AM, Amier R, Ferro DA, Franken L, Nijveldt R, et al: Study design: the effect of aerobic exercise on cerebral perfusion in patients with vascular cognitive impairment. Alzheimers Dement Transl Res Clin Interv 2017;3:157–165.