



Universiteit
Leiden
The Netherlands

Trigger factors and mechanisms in migraine

Schoonman, G.G.

Citation

Schoonman, G. G. (2008, September 11). *Trigger factors and mechanisms in migraine*. Retrieved from <https://hdl.handle.net/1887/13094>

Version: Corrected Publisher's Version

License: [Licence agreement concerning inclusion of doctoral thesis in the Institutional Repository of the University of Leiden](#)

Downloaded from: <https://hdl.handle.net/1887/13094>

Note: To cite this publication please use the final published version (if applicable).

TRIGGER FACTORS AND MECHANISMS IN MIGRAINE

Geurt Gerhard Schoonman (roepnaam: Guus)

Geurt Gerhard Schoonman

Trigger factors and mechanisms in migraine

PhD thesis, Leiden University Medical Center, Leiden 2008

ISBN: 978-90-71382-47-5

Layout by: Gildeprint Drukkerijen B.V., Enschede, The Netherlands

Printed by: Gildeprint Drukkerijen B.V., Enschede, The Netherlands

Cover image courtesy: US Geological Survey/Cascades Volcano Observatory, USA

Copyright of individual chapters lies with the publisher of the journal listed at the beginning of each respective chapter. No part of this thesis may be reproduced in any form, by print, photocopy, digital file, internet or any other means without permission from the author.

The investigations described in this thesis were performed at the department of Neurology of the Leiden University Medical Centre, Leiden, the Netherlands and the department of Neurology of the Zurich University Hospital, Zurich, Switzerland. This work was supported by the Netherlands Organisation for Scientific Research (NWO), grantnumber: 940-38-029

Financial support for the printing of this thesis has been generously provided by: Leiden University, Astra Zeneca B.V., Stichting Het Remmert Adriaan Laan Fonds, Janssen-Cilag B.V., Menarini Farma Nederland, Glaxo Smith Kline, Teva Pharma NL, Sanofi Aventis, Nederlandse Hoofdpijn Vereniging, JE Jurriaanse Stichting

TRIGGER FACTORS AND MECHANISMS IN MIGRAINE

Proefschrift

ter verkrijging van
de graad van Doctor aan de Universiteit Leiden,
op gezag van Rector Magnificus prof.mr. P.F. van der Heijden,
volgens besluit van het College voor Promoties
te verdedigen op donderdag 11 september 2008
klokke 16:15 uur

door

Geurt Gerhard Schoonman
geboren te Deventer in 1974

PROMOTIECOMMISSIE

Promotor: Prof.dr. M.D. Ferrari

Copromotores: Dr. G.M. Terwindt
Dr. J. van der Grond

Referent: Prof.dr. P.R. Saxena

Overige leden Prof.dr. J.G. van Dijk
Prof.dr. M.A. van Buchem

CONTENTS

General introduction and aims of this thesis	7
1. The Prevalence of Premonitory Symptoms in Migraine; A Questionnaire Study in 461 Patients. <i>Cephalalgia 2006; 26: 1209-13.</i>	19
2. Is stress a trigger factor for migraine? <i>Psychoneuroendocrinology 2007;32: 532-538.</i>	27
3. Normobaric hypoxia and nitroglycerin as trigger factors for migraine. <i>Cephalalgia 2006; 26: 816-9.</i>	37
4. Mild Cerebral Edema in Acute Mountain Sickness After Isobaric Hypoxia. A 3 Tesla Magnetic Resonance Imaging Study. <i>Journal of Cerebral Blood Flow and Metabolism. 2008; 28:198-206.</i>	45
5. Magnetic Resonance Angiography of the Human Middle Meningeal Artery: Implications for Migraine. <i>Journal of Magnetic Resonance Imaging. 2006; 24: 918-21.</i>	59
6. Cerebral blood flow response to nitroglycerin predicts the occurrence of a provoked migraine attack <i>Submitted</i>	67
7. Migraine headache is not associated with cerebral or meningeal vasodilatation - a 3T magnetic resonance angiography study. <i>Brain. 2008 May 23 (epub ahead of print)</i>	83
General discussion and conclusions	103
Samenvatting en conclusies	109
References	115
Bibliography	129
Curriculum vitae	131

— |

| —

—————

— |

| —

CHAPTER 1

GENERAL INTRODUCTION



CLINICAL FEATURES OF MIGRAINE

Migraine is a severe paroxysmal neurovascular disorder and considered a major cause of disability by the World Health Organisation^{1,2}. The duration of a migraine attack is between 4 to 72 hours³ and a full blown attack consists of four phases: premonitory, aura, headache and recovery^{4,5}. The premonitory phase can last up to 24 hours and consists of a wide range of symptoms, such as mood disturbances, autonomic symptoms and concentration problems. The prevalence of premonitory symptoms is unclear and ranges from 8%⁶ to 80%⁷ in a clinic based sample. The second phase is the aura phase. Approximately 33% of migraine patients report aura symptoms during an attack⁸ which mostly consist of visual or sensory phenomena⁹. Headache is the third part of an attack and for many patients the most prominent phase. The typical headache during a migraine attack is moderate to severe, unilateral, pounding and aggravates during physical activity. The headache is accompanied by nausea, vomiting and phono/photophobia (Table 1). The final phase of a migraine attack is the recovery phase consisting of symptoms that are similar to the premonitory phase⁷. The clinical presentation of a migraine attack can differ within and between migraine patients⁹.

EPIDEMIOLOGY AND ATTACK SUSCEPTIBILITY

The one year prevalence of migraine in the Netherlands is 25% in women and 7.5% in men⁸ and in the USA the one year prevalence is 17.2% in women and 6% in men¹⁰. Everybody can have a migraine attack, but it is the recurrence of attacks that is abnormal¹¹. A patient is considered a migraine patient only after five MO attacks or two MA attacks according to the IHS criteria³. Attack frequency varies between and within patients and the occurrence of a migraine attack is the result of a misbalance between susceptibility and trigger factors¹². Migraine susceptibility is strongly influenced by genetic factors¹³ and prophylactic treatment¹⁴. Up to now three genes have been identified in familial hemiplegic migraine which is a subtype of migraine with aura¹⁵⁻¹⁷. Whether these genes are involved in the common types of migraine is unknown^{18,19}. Besides genetic factors, prophylactic drugs have shown to alter susceptibility for migraine. Beta-blockers and anti-epileptic drugs are first choice, however, their efficacy is rather limited¹⁴.

Table 1 IHS diagnostic criteria for migraine with and without aura

1.1 Migraine without aura		_____ regel 1
A. At least 5 attacks fulfilling criteria B–D		_____ regel 2
B. Headache attacks lasting 4–72 hours (untreated or unsuccessfully treated)		_____ regel 3
C. Headache has at least two of the following characteristics:	1. unilateral location	_____ regel 4
	2. pulsating quality	_____ regel 5
	3. moderate or severe pain intensity	_____ regel 6
	4. aggravation by or causing avoidance of routine physical activity (eg, walking or	_____ regel 7
	1. nausea and/or vomiting	_____ regel 8
D. During headache at least one of the following:	2. photophobia and phonophobia	_____ regel 9
E. Not attributed to another disorder		_____ regel 10
		_____ regel 11
1.2 Migraine with aura		_____ regel 12
A. At least 2 attacks fulfilling criteria B–D		_____ regel 13
B. Aura consisting of at least one of the following, but no motor weakness:	1. fully reversible visual symptoms including positive features (eg, flickering lights, spots or lines) and/or negative features (ie, loss of vision)	_____ regel 14
	2. fully reversible sensory symptoms including positive features (ie, pins and needles) and/or negative features (ie, numbness)	_____ regel 15
		_____ regel 16
C. At least two of the following:	1. homonymous visual symptoms and/or unilateral sensory symptoms	_____ regel 17
	2. at least one aura symptom develops gradually over ≥ 5 minutes and/or different aura symptoms occur in succession over ≥ 5 minutes	_____ regel 18
		_____ regel 19
D. Headache fulfilling criteria B–D for 1.1 <i>Migraine without aura</i> begins during the aura or follows aura within 60 minutes		_____ regel 20
		_____ regel 21
E. Not attributed to another disorder		_____ regel 22
		_____ regel 23
		_____ regel 24
		_____ regel 25
		_____ regel 26
		_____ regel 27
		_____ regel 28
		_____ regel 29
		_____ regel 30
		_____ regel 31
		_____ regel 32
		_____ regel 33
		_____ regel 34
		_____ regel 35
		_____ regel 36
		_____ regel 37
		_____ regel 38
		_____ regel 39

TRIGGER FACTORS FOR MIGRAINE

A trigger for migraine is any factor that on exposure or withdrawal leads to the development of a migraine attack.²⁰ An extensive list of factors has been proposed as possible trigger factors for migraine (Table 2). Observational questionnaire studies often suggest strong associations between possible trigger factors and migraine which rarely is confirmed by prospective studies and experimental trials. Using questionnaires

regel 1 _____ it is easy to reach a large number of patients, however associations are mainly based
regel 2 _____ on retrospective data and should be regarded as hypothesis generating²¹. On the other
regel 3 _____ hand experimental studies mainly focus on one factor at the time. In the next section,
regel 4 _____ possible trigger factors will be grouped into six categories: food products, stress, female
regel 5 _____ hormones, atmospheric, pharmacological and other factors.

A) Food products

regel 6 _____
regel 7 _____ The occurrence of migraine is often linked to the intake of certain food products and
regel 8 _____ migraine has been described as food allergy²². Despite many studies, the association
regel 9 _____ between food products and migraine remains unclear. Based on retrospective
regel 10 _____ questionnaires a long list possible migraine triggering products has been formulated
regel 11 _____ (Table 2). Among the most frequently mentioned products are alcohol (including
regel 12 _____ wine), cheese, chocolate as well as withdrawal of caffeine and missing a meal.²³⁻³⁰
regel 13 _____ Furthermore, several diet elimination studies suggest a positive association between
regel 14 _____ food and migraine.^{22,31-33} On the other hand, experimental provocation studies are less
regel 15 _____ positive. Red wine provoked migraine in 9 out of 11 migraine patients who were pre-
regel 16 _____ selected on being sensitive for red wine.³⁴ Chocolate triggered migraine in 5 out of
regel 17 _____ 12 "chocolate sensitive" migraine patients³⁵, whereas in a second study the headache
regel 18 _____ response after chocolate did not differ from placebo³⁶. Tyramine 200mg has also been
regel 19 _____ tested in a provocation study in 80 migraine patients and there was no difference in the
regel 20 _____ occurrence of headache between tyramine and placebo.³⁷ Prospective studies in which
regel 21 _____ the intake of food and the occurrence of migraine attacks are scored independently
regel 22 _____ using electronic diaries to prevent retrospective data entries are missing.
regel 23 _____
regel 24 _____

B) Stress

regel 25 _____ Although no clear definition of stress exists³⁸, it has been linked to a whole range
regel 26 _____ of diseases including multiple sclerosis³⁹, asthma⁴⁰ and risk factors for cardiovascular
regel 27 _____ disease.⁴¹ In migraine, both mental and physical stressors are frequently reported as
regel 28 _____ trigger factor. In retrospective questionnaire studies between 30.5% and 81.8% of
regel 29 _____ patients reported psychosocial stressors as trigger factor, whereas between 15.5% and
regel 30 _____ 43.1% of patients identified physical stressors as possible trigger factor (Table 2). Also
regel 31 _____ prospective studies using diaries suggest a positive association between mental stress
regel 32 _____ and migraine.^{42,43} However, this seemingly apparent association between stress and
regel 33 _____ migraine is difficult to replicate in observational and experimental studies using biological
regel 34 _____ stress parameters such as cortisol and cardiovascular parameters. In experimental studies
regel 35 _____ no difference was found in cardiovascular response between the migraine attack and
regel 36 _____ the inter-ictal state.^{44,45}
regel 37 _____
regel 38 _____
regel 39 _____

Table 2 Potential trigger factors for migraine.

	Trigger factor	Response rate* Range (%)	
Food products ^{24-30,119}	Various food items	10 – 36	
	Missing a meal	0.9 – 55.8	
	Chocolate	0 – 22.5	
	Wine	1.4	
	Alcohol	20	
	Dairy products	18.5	
	Caffeine (withdrawal)	6.4	
Atmospheric ^{24-29,120}	Weather changes	6.9 – 52.3	
	Sunlight exposure	4.2 – 38	
	Altitude/ hypoxia		
	Chinook winds		
Stress ^{24-30,121}	Smoking	2 – 26	
	Psychosocial	30.5 – 81.8	
	Physical	15.5 – 43.1	
Female Hormones ^{24-30,122}	Vacation and travel	8 – 54.6	
	Menstruation	20.7 – 53.5	
Pharmacological	Nitroglycerin ^{55,56,78,116,123-129}	20 -83%	
	Sildenafil ⁵⁸	83	
	Dipyridamole ¹³⁰	50	
	Histamine ⁶⁴	50	
	M-chlorophenylpiperazine ⁵⁹	53	
	Calcitonin gene related peptide ⁶¹	33.3	
	Acetazolamide ⁶⁵	Not tested in RCT	
	Prostaglandine E1 ⁶⁸	Not tested in RCT	
	Reserpine ⁶⁹	Not tested in RCT	
	Calcineurin inhibitors ⁷⁰	Not tested in RCT	
	Polidocanol foam ⁷¹	Not tested in RCT	
	Other	Sleep (lack or excess) ⁷²	31 – 52.4
		Visual stimulation ⁷³	
Cerebral angiography ⁷²			
Sexual activity ⁷⁴		0 – 11	
Use of personal computer ²⁴		6.6	

*Response rate are based on findings in questionnaire studies, prospective diary studies or experimental provocation studies.

regel 1 _____
regel 2 _____
regel 3 _____
regel 4 _____
regel 5 _____
regel 6 _____
regel 7 _____
regel 8 _____
regel 9 _____
regel 10 _____
regel 11 _____

C) Female hormones

Based on clinical arguments there is a strong association between female hormones and the occurrence of migraine attacks. The life time prevalence of migraine is 3 times higher in females compared to males⁸, between 20.7% and 53.5% of females reported an association between menstruation and migraine (Table 2) and there is a decrease in migraine frequency during pregnancy.⁴⁶ In a study of 40 female migraine patients, the incidence of migraine attacks was inversely associated with urinary oestrogen concentration across the menstrual cycle. There was no association between migraine and urinary concentrations of progestogens.⁴⁷

regel 12 _____
regel 13 _____
regel 14 _____
regel 15 _____
regel 16 _____
regel 17 _____
regel 18 _____
regel 19 _____
regel 20 _____
regel 21 _____
regel 22 _____
regel 23 _____
regel 24 _____

D) Atmospheric

Weather changes have also been linked to a wide variety of medical diseases⁴⁸ including migraine.⁴⁹ Retrospective questionnaires showed that between 6,9% and 52,3% of migraine patients identify weather changes as possible trigger factor (Table 2). In contrast three prospective studies, combining objective weather data from meteo institutes with information from headache diaries or visits to the emergency room for migraine, showed no positive associations.⁵⁰⁻⁵² Only one study found a positive relation between weather changes and the occurrence of headache in 77 migraine patients.⁵³ Furthermore there is a large discrepancy between what patients think and what can be objectified. For instance a positive association between Chinook winds and migraine attacks was suggested by 88% of 34 migraine patients, whereas an objective correlation could only be found in 21% of patients.⁵⁴ Experimental studies including atmospheric parameters are limited in number.

regel 25 _____
regel 26 _____
regel 27 _____
regel 28 _____
regel 29 _____
regel 30 _____
regel 31 _____
regel 32 _____
regel 33 _____
regel 34 _____
regel 35 _____
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

E) Pharmacological

Nitroglycerin (NTG) is frequently used in migraine provocation studies (Table 2). The clinical response after NTG (0.5 micrograms/kg/20min) consists of an immediate type headache during infusion and a delayed headache attack after 5 to 6 hours which fulfils the criteria of migraine without aura in 20% to 83% of patients (Table 2). Migraine patients without aura might be more susceptible to nitroglycerin than patients with aura.^{55,56} Sildenafil (Viagra) is a highly selective phosphodiesterase type 5 inhibitor used to treat patients with erectile dysfunction⁵⁷ and in migraine susceptible patients Viagra has shown to provoke delayed migraine attacks in 10 out of 12 patients.⁵⁸ A third drug shown to provoke migraine is m-chlorophenylpiperazine (mCPP). Migraine attacks were triggered in 10 out of 19 migraine patients (53%) in a randomized controlled trial.⁵⁹ Also in a study including patients with bulimia and anorexia nervosa, mCPP triggered severe headache 28 out of 52 patients (54%).⁶⁰ Calcitonin gene related

peptide (CGRP) is a vasoactive peptide that is increased during spontaneous migraine attacks⁶¹. In turn, infusion of CGRP triggers migraine in 3 out of 9 susceptible migraine patients.⁶² The neurotransmitter histamine has also shown to trigger moderate to severe throbbing headache in migraine susceptible patients⁶³ fulfilling the criteria for migraine in 50% of the migraine patients.⁶⁴ Besides aforementioned drugs, several others drugs might be capable of triggering migraine, but they are up to now never been tested in a formal randomized controlled trial (RCT). Acetazolamide (Diamox), a carbonic anhydrase inhibitor, is both used to provoke and to treat migraine. Oral administration of acetazolamide (14.3 mg/kg) in 20 migraine patients caused migraine headache accompanied by photophobia, phonophobia and nausea after 1 to 8 hours.⁶⁵ The number of patients fulfilling the criteria for migraine was not specified in this study. In contrast, diamox (500 to 750 mg daily) has also been used as treatment in migraine and it might be effective in the acute treatment of migraine aura status.⁶⁶ Furthermore, diamox (500mg) has been tested as prophylaxis for migraine in 53 patients and was not effective.⁶⁷ And finally prostaglandine E1⁶⁸, reserpine⁶⁹, calcineurin inhibitors (eg, cyclosporine and tacrolimus) ⁷⁰ and polidocanol foam⁷¹ might be able to provoke migraine attacks in susceptible patients.

F) Other possible trigger factors

Sleep (lack or excess) and fatigue are frequently associated with migraine attacks (Table 2). Also in a prospective diary study the quality of sleep seemed to be negatively associated with the occurrence of migraine attacks.⁷² Visual stimulation has been used to trigger migraine in a fMRI study.⁷³ Two (out of 10) migraine patients with aura experienced a typical migraine aura and 8 (out of 12) experienced migraine headache within 7.3 minutes after provocation. Whether these headache episodes fulfilled migraine criteria was not described. Cerebral angiography using contrast agent has shown to induce headache in 15 (out of 45) patients after 2 hours.⁷² In four patients (8.8%) symptoms fulfilled criteria for migraine without aura. Sexual activity has also been associated with a wide range of positive as well as negative effects, including headache and migraine (Table 1). There is even a sub classification for "preorgasmic" and "orgasmic" headache.³ In a group of 51 patients with "headache associated with sexual activity" co morbidity with migraine was 25%.⁷⁴ Whether it is just physical stress causing headache or something extra during sexual activity is not known. The use of personal computer (PC) is a rather new factor and identified as possible trigger factor in 6.6% of Japanese migraine patients.²⁴ This factor has not been included in other questionnaire studies or experimental trials.

PATHOPHYSIOLOGY OF A SPONTANEOUS MIGRAINE ATTACK

Activation of the trigeminovascular system is pivotal during the headache phase of a migraine attack⁷⁵. The mechanism causing activation of the trigeminovascular system remains to be elucidated^{12,76}. Several mechanisms might be involved in the initiation of a migraine attack. A) Cortical spreading depression (CSD) is a steady depolarization of neuroglial membranes and is the pathophysiological mechanism underlying migraine aura¹². A long-lasting blood flow change in meningeal arteries have been observed after CSD depending on trigeminal and parasympathetic activation⁷⁷. B) Vasodilatation of cerebral and meningeal arteries might activate trigeminal nerves. Vasoactive substances such as nitroglycerin can trigger migraine in susceptible patients⁷⁸ and triptans may exert their anti-migraine effect through vasoconstriction of cranial blood vessels⁷⁵. C) Neurogenic inflammation caused by vasoactive peptides released from the trigeminal nerve or other sources such as blood have shown to activate and sensitize meningeal perivascular nerve ending causing activation of the trigeminovascular system⁷⁹ and possible disruption of the blood-brain barrier⁸⁰. D) Nociceptive information from the trigeminal nerve is modulated in the brainstem⁸¹. Activation of brainstem area's, such as the peri-aqueductal grey, has been shown during spontaneous and provoked migraine attacks^{82,83}. E) The occurrence of premonitory symptoms (such as fluid retention, sleep problems and food craving) prior to the onset of headache suggest involvement of the hypothalamus.^{84,85} Hypothalamic activation has also been shown in other trigeminal neuralgias, such as cluster headache⁸⁶. For further information on the pathophysiology of migraine please read some excellent reviews that have been published recently^{12,75,79,87}.

MECHANISM OF ACTION OF TRIGGER FACTOR IN MIGRAINE: STRESS, HYPOXIA AND NITROGLYCERIN

As presented, there are many (potential) trigger factors for migraine all with a different mechanism of action. Since it is not feasible to study all we will focus on three trigger factors: mental stress, normobaric hypoxia and nitroglycerin. The study of trigger factor mechanisms may provide further insight into the first phases of a migraine attack

A) Stress and the autonomic nervous system during migraine

Mental stressors are commonly perceived as important trigger factors by both patients and physicians⁸⁸, although direct evidence for this claim is lacking. In retrospective questionnaire studies, up to 62% of migraine patients reported that psychosocial stress

was an important trigger-factor for their attacks^{25,29,89}, but patients have a tendency to overestimate stress on retrospective measures⁹⁰. In cross-sectional studies, migraine patients were found to have elevated plasma levels of cortisol, an indicator for stress, both outside a migraine attack compared to healthy volunteers⁹¹ and during attacks compared to the inter-ictal phase⁹². Stress-provocation studies, involving mental and physical stressors, have suggested sympathetic and parasympathetic changes in migraine patients outside attacks compared to healthy volunteers⁹³⁻⁹⁶. However, experimental prospective studies examining whether stress-related biological changes are actually temporally related to the onset of migraine attacks, are lacking. We therefore performed a prospective, longitudinal ambulatory study, assessing perceived stress and objective stress-related biological changes in the four days prior to an impending migraine attack (chapter 2).

B) Hypoxia and blood brain barrier dysfunction

Hypoxia might also be a trigger factor for migraine. Firstly, acute exposure to high altitude may induce acute mountain sickness (AMS), which is characterized by headache, insomnia, dizziness, lassitude, fatigue and gastrointestinal symptoms such as anorexia, nausea, or vomiting in an unacclimatized person who has recently reached an altitude above 2500 m⁹⁷. Up to one third of subjects with acute AMS also fulfill the criteria for migraine^{3,98,99}. Secondly, chronic exposure to high altitude is associated with an increased migraine prevalence^{100,101} and thirdly, sumatriptan is an established drug for the acute treatment of migraine⁷⁵, and was also shown to be effective in some studies in AMS^{102,103}. In chapter 3 we have tested whether normobaric hypoxia may trigger migraine attacks in migraine patients under experimental conditions. Hypoxia has many biological effects and one of the mechanisms involved in the pathophysiology of AMS is disruption of the BBB causing cerebral edema⁹⁷. In severe cases of AMS there are clear signs of vasogenic edema as shown by MRI¹⁰⁴. Also in migraine disruption of the BBB has been suggested¹⁰⁵. Whether hypoxia causes cerebral edema in mild cases of AMS (resembling migraine) is unclear. This question was studied in chapter 4.

C) Nitroglycerin and changes in cerebral blood flow

Nitroglycerin is an exogenous donor of nitric oxide¹⁰⁶, which is involved in central pain mechanism¹⁰⁷ and regulation of cerebral blood flow¹⁰⁸. Infusion of NTG has shown to increase the diameter of the middle cerebral artery¹⁰⁹ and meningeal media artery¹¹⁰ as well as to decrease blood flow velocity in the internal carotid artery and middle cerebral artery¹¹¹⁻¹¹³. The effects of NTG on cerebral blood flow are caused either through the release of CGRP from the trigeminal nerve^{114,115} or via a direct effect on vascular smooth

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____ muscle cells in blood vessels¹⁰⁶. Infusion of NTG results in immediate type headache
regel 2 _____ in >80% of migraine patients and <20% in healthy volunteers¹¹⁶. A delayed migraine
regel 3 _____ attack is observed several hours after infusion of NTG in approximately 60% to 80%
regel 4 _____ of migraine patients and very rarely in healthy volunteers without a family history of
regel 5 _____ migraine^{55,78,116}. Whether there is a difference in cerebrovascular response to NTG
regel 6 _____ between migraine patients and healthy controls is unclear. One study suggested an
regel 7 _____ increased cerebrovascular response during NTG infusion in migraine patients¹¹⁷, whereas
regel 8 _____ in a second study no increased response was observed.¹¹⁸ This will be studied in chapter
regel 9 _____ 6. In the same provocation study (chapter 7) we have studied cerebrovascular changes
regel 10 _____ (both blood vessel diameters and blood flow) during the provoked migraine attack.

AIMS OF THIS THESIS

regel 11 _____
regel 12 _____
regel 13 _____
regel 14 _____
regel 15 _____ As discussed there are many potential trigger factors for migraine. We have chosen
regel 16 _____ to study three (potential) trigger factors: mental stress, normobaric hypoxia and
regel 17 _____ nitroglycerin. The following aims for this thesis were defined:

- regel 18 _____
regel 19 _____ 1. To assess the prevalence of premonitory symptoms in a clinic based sample of migraine
regel 20 _____ patients and to study a potential overlap between premonitory symptoms and trigger
regel 21 _____ factors (chapter 1).
- regel 22 _____
regel 23 _____ 2. To assess both subjective and objective stress related parameters during the
regel 24 _____ development of a spontaneous migraine attack (chapter 2).
- regel 25 _____
regel 26 _____ 3. To test normobaric hypoxia as a trigger factor for migraine in migraine susceptible
regel 27 _____ patients and to compare the response to nitroglycerin (chapter 3).
- regel 28 _____
regel 29 _____ 4. To test whether normobaric hypoxia caused cerebral edema in healthy volunteers
regel 30 _____ (chapter 4).
- regel 31 _____
regel 32 _____ 5. To develop a method to measure vasodilatation in cranial blood vessels as small as
regel 33 _____ the middle meningeal artery in healthy volunteers and migraine patients using magnetic
regel 34 _____ resonance angiography (chapters 5 and 6).
- regel 35 _____
regel 36 _____ 6. To assess the initial vascular response to nitroglycerin in migraine as a predictor for
regel 37 _____ the development of a provoked migraine attack (chapter 6).
- regel 38 _____
regel 39 _____

7. To assess vasodilatation in cranial blood vessels during a provoked migraine attack (chapter 7).

- ___ regel 1
- ___ regel 2
- ___ regel 3
- ___ regel 4
- ___ regel 5
- ___ regel 6
- ___ regel 7
- ___ regel 8
- ___ regel 9
- ___ regel 10
- ___ regel 11
- ___ regel 12
- ___ regel 13
- ___ regel 14
- ___ regel 15
- ___ regel 16
- ___ regel 17
- ___ regel 18
- ___ regel 19
- ___ regel 20
- ___ regel 21
- ___ regel 22
- ___ regel 23
- ___ regel 24
- ___ regel 25
- ___ regel 26
- ___ regel 27
- ___ regel 28
- ___ regel 29
- ___ regel 30
- ___ regel 31
- ___ regel 32
- ___ regel 33
- ___ regel 34
- ___ regel 35
- ___ regel 36
- ___ regel 37
- ___ regel 38
- ___ regel 39

— |

| —

—————

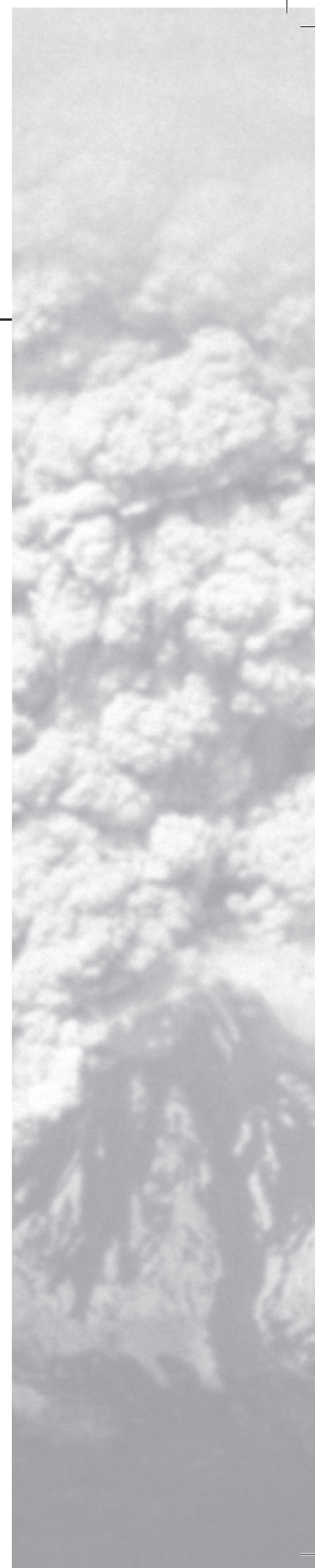
— |

| —

CHAPTER 1

THE PREVALENCE OF PREMONITORY SYMPTOMS IN MIGRAINE: A QUESTIONNAIRE STUDY IN 461 PATIENTS

Cephalalgia 2006;26:1209-13



ABSTRACT

Migraine attacks are often preceded by premonitory symptoms. Prevalence rates of migraine patients reporting one or more premonitory symptoms show considerable variability and rates range between 12% and 79%. Sources of variability might be differences in study population or research design. Using a questionnaire we retrospectively studied the prevalence of 12 predefined premonitory symptoms in a clinic based population. Of 461 migraine patients, 374 responded (81%). At least one premonitory symptom was reported by 86.9%, and 71.1% reported two or more. The most frequently reported premonitory symptoms were fatigue (46.5%), phonophobia (36.4%) and yawning (35.8%). The mean number of premonitory symptoms per person was 3.2 (\pm 2.5). Women reported 3.3 premonitory symptoms compared to 2.5 symptoms in men ($p=0.01$). Age, education, migraine subtype (with or without aura), and mean attack frequency had no effect on the mean number of symptoms per individual. In conclusion, premonitory symptoms are frequently reported by migraine patients. Sensitivity and specificity of premonitory symptoms for migraine need to be assessed using prospective methods.

INTRODUCTION

Migraine is a severe paroxysmal neurovascular disorder and considered a major cause of disability by the World Health Organization¹. The primary cause of a migraine attack is unknown but probably lies within the central nervous system². Prior to the start of the headache phase several non-headache symptoms (often called premonitory symptoms) are reported by migraine patients, such as changes in mood, behavior and sensory perception⁴. In a selected population migraine patients were able to predict an upcoming migraine attack well before the start of the headache phase¹³¹. Prevalence rates of patients reporting one or more premonitory symptoms ranges between 12%¹³² and 79%¹³³. One source of variability in prevalence rate might be differences in study population. In population based studies rates range from 12% in migraine patients without aura to 18% in migraine patients with aura¹³², whereas in clinic based studies prevalence rates range from 33%^{134,135} to 79%¹³³. Other sources of variability might be differences in study design such as preselection of patients or unclear definitions of premonitory symptoms. In this study we assessed the prevalence of 12 frequently reported premonitory symptoms using a questionnaire in a large unselected clinic based population and only symptoms preceding 2/3 of attacks or more were considered a premonitory symptom.

METHODS

Migraine patients (diagnosed according to the criteria of the IHS³) from the Neurology outpatient clinic of the Leiden University Medical Centre received a questionnaire by mail. A reminder was send out to the patients who had not responded after 8 weeks. The questionnaire addressed migraine characteristics, sociodemographic factors and possible premonitory symptoms. Migraine related variables were: migraine subtype (migraine with or without aura according to the criteria of IHS³) and mean attack frequency per month in the last half year. The following sociodemographic variables were included: age, sex and education in 3 categories: primary school or low vocational training, middle academic/vocational training, and higher academic/vocational training. Twelve possible premonitory symptoms were included based on reports in the literature^{4,131,135}: Concentration problems, depression, food craving, physical hyperactivity, irritability, nausea, phonophobia, fatigue, sleep problems, stressed feeling, stiff neck and yawning. For every possible premonitory symptom patients answered the question: "How often is a migraine attack preceded by this symptom?" Answers were categorized as never, less

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

than 1/3 of attacks, 1/3 to 2/3 of attacks or in more than 2/3 of attacks. Photophobia was not included in the questionnaire since co-occurrence of aura symptoms and visual hypersensitivity might introduce bias. The duration of the premonitory phase was not strictly defined. The local ethical committee had approved the study. Symptoms were considered a premonitory symptom when at least 2/3 of migraine attacks were preceded by this particular symptom.

Prevalence of every premonitory symptom was calculated and presented as percentage. The number of premonitory symptoms per individual was calculated and presented as mean (and SD). A difference in mean number of symptoms between subgroups was tested using the non-paired t-test (for sex and migraine subtype) or one-way ANOVA (for age, education and attack frequency). In case of non-normality the Mann-Whitney U test or Kruskal Wallis test were used. The Bonferroni correction was applied for multiple testing and a p value <0.01 was considered significant. The co-occurrence of PS within patients was tested using Spearman's rank correlation coefficient and presented as correlation matrix.

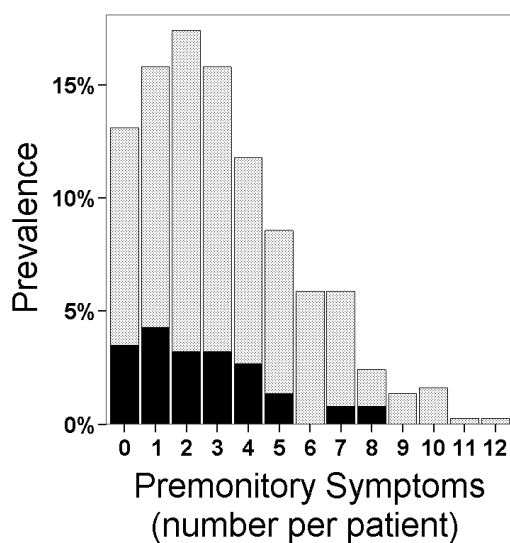


Figure 1 Number of premonitory symptoms per subject. Black bars represent males, gray bars females.

RESULTS

The questionnaire was sent to 461 migraine patients; 374 (81%) responded. The characteristics of the study population are shown in Table 1. Forty-nine patients (13.1%)

reported no premonitory symptoms, 86.9% of patients reported at least one symptom and 71.1% reported two or more (Figure 1). The most frequently reported premonitory symptoms were fatigue (46.5%), phonophobia (36.4%) and yawning (35.8%) (Table 2). The mean number of premonitory symptoms reported per person was 3.2 (SD 2.5). Women reported a mean of 3.3 symptoms compared to a mean of 2.5 in men ($p=0.01$). The effects of age, education, migraine subtype, and mean attack frequency on the mean number of symptoms per individual were not statistically significant (Table 1). Of the migraine patients 52% had migraine with aura (Table 1). No significant difference in premonitory symptoms was found between migraine subtypes (with and without aura) (Table 2). The co-occurrence of symptoms is presented in Table 3. Depression and irritability showed the strongest correlation, followed by depression and concentration problems and depression and a stressed feeling.

Table 1 Migraine and sociodemographic properties of all interviewed patients.

	Subgroups	N (%)	Mean number of PS per individual (SD)	
Total population		374	3.2 (2.5)	
Sex				
	Male	74 (20%)	2.5 (2.1)	
	Female	300 (80%)	3.3 (2.5)	p=0.01
Age (years)				
	<30	29 (8%)	3.6 (2.5)	
	30-50	172 (46%)	3.0 (2.2)	
	50>	173 (46%)	3.2 (2.7)	p=0.59
Education				
	low	147 (39%)	3.5 (2.4)	
	middle	78 (21%)	2.9 (2.7)	
	high	148 (39%)	3.0 (2.5)	p=0.03
Migraine subtype				
	without aura	179 (48%)	2.9 (2.4)	
	with aura	195 (52%)	3.4 (2.6)	p=0.12
Attack frequency (per month)				
	<2	94 (25%)	2.9 (2.4)	
	2-4	139 (37%)	3.1 (2.4)	
	>4	140 (38%)	3.3 (2.6)	p=0.65

Table 2 Prevalence of premonitory symptoms

Premonitory symptom	Prevalence (%)			P value	MO	MA	P value
	All patients (N=374)	Male (N=74)	Female (N=300)				
Fatigue	46.5	39.1	48.3	0.16	47.5	45.6	0.72
Phonophobia	36.4	24.3	39.3	0.02	30.7	41.5	0.03
Yawning	35.8	31.1	37.0	0.34	34.6	36.9	0.65
Stiff neck	35.0	32.4	35.7	0.60	40.8	29.7	0.03
Nausea	28.6	16.2	31.7	0.008	22.9	33.8	0.02
Concentration problems	28.1	29.7	27.8	0.74	20.7	35.1	0.002
Irritability	28.1	25.6	28.6	0.59	24.0	32.0	0.09
Depression	17.6	13.5	18.6	0.29	18.4	16.9	0.70
Craving	17.4	6.7	20.0	0.007	14.0	20.5	0.10
Stressed feeling	15.2	14.8	15.3	0.92	14.0	16.4	0.51
Physical hyperactivity	15.0	6.7	17.0	0.03	12.8	16.9	0.27
Sleep problems	13.9	10.8	14.6	0.39	14.0	13.9	0.98

*Prevalence is the percentage of patients of the total population (or subgroup) reporting a certain symptoms. MO denotes migraine without aura, MA migraine with aura.

Table 3 Co-occurrence of premonitory symptoms: Spearman's rank correlation coefficient matrix. Field shading indicates correlation strength.

	SF	SN	PHH	IR	YA	DE	FA	CR	PH	CP	NA	SP
Stressed feeling (SF)												
Stiff neck (SN)	,234											
Physical hyperactivity	,197	,116										
Irritability (IR)	,198	,126	,171									
Yawning (YA)	-,038	,129	,171	,144								
Depression (DE)	,350	,160	,179	,397	,151							
Fatigue (FA)	,171	,203	,149	,290	,220	,313						
Craving (CR)	,120	,048	,262	,200	,113	,084	,053					
Phonophobia (PH)	,082	,144	,228	,306	,084	,190	,164	,211				
Concentration problems (CP)	,132	,101	,137	,324	,057	,350	,267	,137	,294			
Nausea (NA)	,044	,130	,049	,130	,206	,188	,181	,100	,186	,170		
Sleep problems (SP)	,109	,125	,004	,127	,024	,138	,153	,121	,194	,075	,104	

DISCUSSION

The proportion of migraine patients reporting premonitory symptom was high: 86.9% of patients reported at least one symptom. This high prevalence rate is comparable to one previous clinic based study where the rate was 79%¹³³, but in contrast with two other studies where rates were about 33%^{134,135}. Variability in rates might be explained by differences in study design such as preselection of patients¹³³ or differences in symptoms that are included in the questionnaire¹³⁵. Furthermore, the study of Amery¹³³ was conducted before the introduction of the IHS migraine criteria. Another source of variability might be the studied population. For instance prevalence rates in population based studies have shown to be as low as 12%¹³². It may be that patients identified in a population based setting are not informed about premonitory symptoms in migraine and, therefore, are less aware of these symptoms. Fatigue was the most common premonitory symptom and the order of reported symptoms is comparable with a previous study in a selected population¹³¹. In our study the percentage of patients presenting with aura was high. Patients with aura are more likely to consult a neurologist than patients without aura and this differences might be increased due to the fact that all patients in the Netherlands see there General Practioner first in case of complaints. However, no significant difference in PS was seen between migraine subtypes.

Females reported more premonitory symptoms than males. An overlap between premonitory symptoms and premenstrual syndrome might explain this difference¹³⁶. Furthermore more females reported craving and nausea as premonitory symptom compared to males. This is an interesting finding since chocolate and sweet cravings are more common in females than males¹³⁷. Nausea is also more frequently reported in females than in males in acute myocardial infarction¹³⁸ and after anaesthesia¹³⁹. The physiological basis for this gender difference is not clear. Besides gender differences co-occurrence of premonitory symptoms within one subject were studied. The strongest associations were found between depression and symptoms such as irritability, concentration problems and fatigue. Co-occurrence of these mood symptoms might not be a coincidence since they are all part of the DSM IV criteria for dysthymic disorder and major depression¹⁴⁰.

There might also be an overlap between premonitory symptoms and trigger factors in migraine. A migraine trigger is any factor that on exposure or withdrawal leads to the development of a migraine attack whereas PS are a consequence of an ongoing attack. For instance mental stress (either the acute episode or the relieve period after an acute episode) is often considered a trigger factor in retrospective questionnaires. However, it is unclear whether migraine attacks can be triggered in an experimental provocation

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____ study¹⁴¹. So, It could be that mental stress trigger a migraine attack or that patients
regel 2 _____ perceive more mental stress because they are in the premonitory phase of a migraine
regel 3 _____ attack. Future prospective diary studies or experimental studies are needed to address
regel 4 _____ this question.

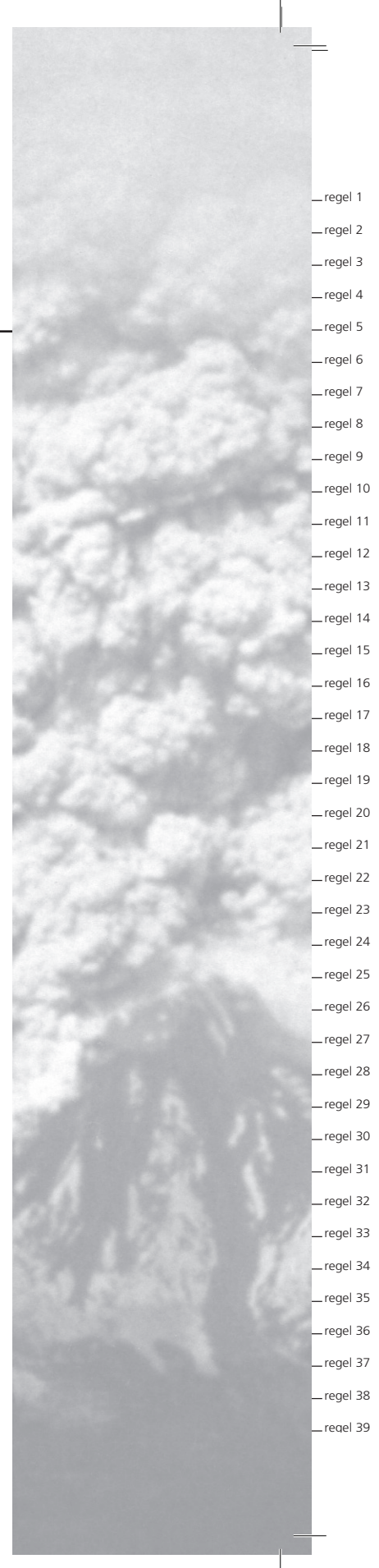
regel 5 _____ This study, as well as other retrospective studies assessing premonitory symptoms in
regel 6 _____ migraine, has some limitations. First, the list of possible premonitory symptoms is based
regel 7 _____ on previous studies^{4,131,135} and may seem somewhat arbitrary. To be complete one should
regel 8 _____ do a full exploration of all possible symptoms associated with a migraine attack. Second,
regel 9 _____ non-responders might have introduced some bias. However, the response rate was 81%
regel 10 _____ and there was no difference in age, sex or migraine subtype between responders and
regel 11 _____ non-responders (data not shown). Third, when should a symptom be classified as a
regel 12 _____ premonitory symptom? We excluded photophobia as a premonitory symptom but it
regel 13 _____ could be argued that phonophobia and nausea are actually part of the headache phase
regel 14 _____ and therefore no PS. Furthermore, in this study we considered symptoms as premonitory
regel 15 _____ symptom if 2/3 of attacks were preceded by this particular symptom. In order to assess
regel 16 _____ sensitivity and specificity of individual premonitory symptoms for migraine attacks,
regel 17 _____ possible premonitory symptoms and migraine attacks need to be studied prospectively
regel 18 _____ preferably^{131,142}. Also the temporal relation between possible premonitory symptoms,
regel 19 _____ aura and the occurrence of headache needs to be assessed in a prospective design.

regel 20 _____ In conclusion, premonitory symptoms are frequently reported by migraine patients.
regel 21 _____ Sensitivity and specificity of premonitory symptoms for migraine need to be assessed
regel 22 _____ using prospective methods.

CHAPTER 2

IS STRESS A TRIGGER FACTOR FOR MIGRAINE?

Psychoneuroendocrinology 2007;32:532-8



— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

ABSTRACT

Background

Although mental stress is commonly considered to be an important trigger factor for migraine, experimental evidence for this belief is lacking.

Objective: To study the temporal relationship between changes in stress related parameters (both subjective and objective) and the onset of a migraine attack.

Methods

This was a prospective, ambulatory study in 17 migraine patients. We assessed changes in perceived stress and objective biological measures for stress (saliva cortisol, heart rate average [HRA], and heart rate variability [low frequency power and high frequency power]) over four days prior to the onset of spontaneous migraine attacks. Analyses were repeated for subgroups of patients according to whether or not they felt their migraine to be triggered by stress.

Results

There were no significant temporal changes over time for the whole group in perceived stress ($p=0.50$), morning cortisol ($p=0.73$), evening cortisol ($p=0.55$), HRA ($p=0.83$), low frequency power ($p=0.99$) and high frequency power ($p=0.97$) prior to or during an attack. Post-hoc analysis of the subgroup of nine stress-sensitive patients who felt that $>2/3$ of their migraine attacks were triggered by psychosocial stress, revealed an increase for perceived stress ($p=0.04$) but no changes in objective stress response measures. At baseline this group also showed higher scores on the Penn State Worry Questionnaire ($p=0.003$) and the Cohen Perceived Stress Scale ($p=0.001$) compared to non stress-sensitive patients.

Conclusions

Although stress-sensitive patients, in contrast to non stress-sensitive patients, may perceive more stress in the days before an impending migraine attack, we failed to detect any objective evidence for a biological stress response before or during migraine attacks.

INTRODUCTION

Migraine is a multifactorial brain disorder characterised by recurrent, disabling attacks of headache, associated autonomic features and, in one third of patients, neurological aura symptoms⁷⁵. Although the pathogenesis of the migraine features is reasonably well understood, it is not clear how migraine attacks are actually triggered. Mental stressors are psychological events that in potential threaten homeostasis of a living organism¹⁴³ and they are commonly perceived as important trigger factors by both patients and physicians⁸⁸, although direct evidence for this claim is lacking. In retrospective questionnaire studies, up to 62% of migraine patients reported that psychosocial stress was an important trigger-factor for their attacks^{25,29,89}, but patients have a tendency to overestimate stress on retrospective measures⁹⁰. In cross-sectional studies, migraine patients were found to have elevated plasma levels of cortisol, an indicator for stress, both outside a migraine attack compared to healthy volunteers⁹¹ and during attacks compared to the inter-ictal phase⁹². Stress-provocation studies, involving mental and physical stressors, have suggested sympathetic and parasympathetic changes in migraine patients outside attacks compared to healthy volunteers⁹³⁻⁹⁶. However, experimental prospective studies examining whether stress-related biological changes are actually temporally related to the onset of migraine attacks, are conspicuously lacking. We therefore performed a prospective, longitudinal ambulatory study, assessing perceived stress and objective stress-related biological changes in the four days prior to an impending migraine attack. We included both patients who claimed that stress would trigger the majority of their attacks (stress-sensitive) and patients who denied such a relationship (non stress-sensitive).

METHODS

Subjects

A total of 69 migraine patients were recruited from our headache outpatient clinic and 27 patients were included in the study. Inclusion criteria were (1) diagnosis of migraine with or without aura according to the criteria of the IHS (code 1.1. and 1.2.1;³ and at least one migraine attack per month in the previous six months. Exclusion criteria were (1) pure menstrual migraine, (2) more than 15 days of headache per month, (3) use of beta-blockers and (4) inability to differentiate between migraine and other types of primary headache syndromes. We asked the patients whether they felt that their attacks were triggered by stress and if so, in what proportion. Patients who claimed that

___ regel 1

___ regel 2

___ regel 3

___ regel 4

___ regel 5

___ regel 6

___ regel 7

___ regel 8

___ regel 9

___ regel 10

___ regel 11

___ regel 12

___ regel 13

___ regel 14

___ regel 15

___ regel 16

___ regel 17

___ regel 18

___ regel 19

___ regel 20

___ regel 21

___ regel 22

___ regel 23

___ regel 24

___ regel 25

___ regel 26

___ regel 27

___ regel 28

___ regel 29

___ regel 30

___ regel 31

___ regel 32

___ regel 33

___ regel 34

___ regel 35

___ regel 36

___ regel 37

___ regel 38

___ regel 39

regel 1 _____ >2/3 of their attacks were triggered by stress were considered “stress sensitive” and
regel 2 _____ those who reported that <2/3 of their attacks was triggered by stress were considered
regel 3 _____ “stress non-sensitive”. The study was approved by the local medical ethical committee
regel 4 _____ and the subjects gave informed consent prior to the start of the study. The study was
regel 5 _____ conducted in the period January to August 2004.

Procedure

regel 6 _____ Patients filled out two stress questionnaires at the start of the observation period.
regel 7 _____ The first was the Cohen Perceived Stress Scale (Cohen PSS)¹⁴⁴ which is a measure for
regel 8 _____ perceived stress in the past month. It is a 14 item questionnaire and the score ranges
regel 9 _____ from 0 (no stress) to 56 (maximum stress). The second questionnaire was the Penn State
regel 10 _____ Worry Questionnaire (PSWQ)¹⁴⁵, a 16 item questionnaire to assess the trait of worrying
regel 11 _____ (ranging from 16 (minimal worries) to 80 (maximum)). Both questionnaires are used to
regel 12 _____ characterize the study population.
regel 13 _____

regel 14 _____ The observation period started at least three days after an attack and lasted up to the
regel 15 _____ first day of the next attack. Migraine symptoms and stress events were scored daily
regel 16 _____ around 22.00 hours using an electronic diary (described below). Saliva samples were
regel 17 _____ taken 3 times per day (30 and 45 minutes after waking up and around 22.00 hours,
regel 18 _____ before filling out the stress and migraine questionnaire); Heart rate was measured daily
regel 19 _____ between 18.00 and 22.00 hours using an ambulatory monitoring system. The timings
regel 20 _____ were chosen in such a way that the recordings would be influenced as little as possible
regel 21 _____ by physical activity during the day.
regel 22 _____

Perceived daily stress and migraine symptoms

regel 23 _____ ‘Personal digital assistants’ devices (Palm Tungsten E) were used as electronic diaries.
regel 24 _____ Data were entered daily around 22.00 hours using a database application (Pendragon
regel 25 _____ Forms 3.2, Pendragon Software Corporation, Libertyville, USA)¹⁴⁶. Perceived daily stress
regel 26 _____ was measured with the validated Daily Stress Inventory (DSI). In short, this is a 58 item
regel 27 _____ inventory of events experienced in the last 24 hours¹⁴⁷. The amount of stress felt in
regel 28 _____ response to each event is rated on a Likert-type scale (0 = event did not happen, 1 =
regel 29 _____ event occurred but was not stressful to 7 = event caused panic). The perceived daily
regel 30 _____ stress is the sum total of all ratings (DSI-sum). Migraine symptoms were assessed using
regel 31 _____ the criteria of the IHS. The diaries were easy to use and retrospective data entry or
regel 32 _____ alterations were disallowed by the PDA program. An alarm sounded daily at 22.00
regel 33 _____ hours to remind patients to fill out the questionnaires.
regel 34 _____
regel 35 _____
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

Salivary cortisol

Saliva samples for cortisol assessment were obtained with 'Salivette' saliva collection tubes (Sarstedt, Germany). Each day patients collected three saliva samples. 30 minutes and 45 minutes after waking up and around 22.00 hours. Patients were instructed not to eat, exercise, smoke or brush their teeth 30 minutes prior to sampling. Patients stored the samples at 7 °C until the end of the observation period. At the end of the observation period patients were asked to report sampling problems. After centrifugation, samples were stored at -80 °C until analysis. Cortisol concentrations were determined using Elecsys 2010 (Roche Diagnostics, Mannheim, Germany). The functional sensitivity of this assay is 2 nmol/L¹⁴⁸.

Heart rate and heart rate variability

Heart rate was measured using the Vrije Universiteit Ambulatory Monitoring System (VU-AMS, version 4.6, Vrije Universiteit Amsterdam)¹⁴⁹ between 18.00 and 22.00 hours during periods of 10 minutes every half hour. R-R wave intervals were recorded on line from a 3-lead ECG. Fast Fourier transformation was used to calculate spectral power of the RR interval¹⁵⁰; a trend was removed from the data to reduce the influence of very low frequencies. A cubic spline function corrected for missing values in the time series to result in regularly sampled time series. The data were multiplied by a Tukey window and transformed from the time domain to the frequency domain with the discrete Fourier transform. The spectra were smoothed by a triangular window (width ~0.01 cycles per RR interval). After integration of the area under the curve, the low frequency (0.05-0.15 Hz) power (LF), reflecting a mix of sympathetic and parasympathetic activity, and the high frequency (0.15-0.30Hz) power (HF), largely reflecting parasympathetic activity were calculated.

Statistical analysis

Temporal changes and differences between the two stress sensitive subgroups in perceived stress, cortisol (morning and evening), HRA, LF and HF power were analysed using a linear mixed model, with observation day and subgroup as fixed factors. A maximum of four pre-migraine days were included in the analysis since the premonitory phase may start up to 48 hours prior to the onset of the headache phase^{4,134}. Cohen PSS and PWSQ differences between stress sensitive subgroups were tested using an unpaired t-test. The Bonferroni correction was applied for multiple testing and P<0.025 was considered significant.

— regel 1
 — regel 2
 — regel 3
 — regel 4
 — regel 5
 — regel 6
 — regel 7
 — regel 8
 — regel 9
 — regel 10
 — regel 11
 — regel 12
 — regel 13
 — regel 14
 — regel 15
 — regel 16
 — regel 17
 — regel 18
 — regel 19
 — regel 20
 — regel 21
 — regel 22
 — regel 23
 — regel 24
 — regel 25
 — regel 26
 — regel 27
 — regel 28
 — regel 29
 — regel 30
 — regel 31
 — regel 32
 — regel 33
 — regel 34
 — regel 35
 — regel 36
 — regel 37
 — regel 38
 — regel 39

RESULTS

Study population and observation periods

Of the 27 patients included in the study, 17 patients had a migraine attack during the observation period (Table 1). In 10 patients we did not measure an attack: six patients dropped out because the ambulatory cardiovascular measurements interfered too much with daily activities and four patients did not have a migraine attack within the observation period. The duration of the pre-ictal observation period in the 17 patients who had a migraine attack was four days in 12 patients, three days in two patients, two days in two and only 1 day in one patient. Some patients developed an attack within a few days after starting the observation period which is the reason for the variability in observation duration. In 12 patients the migraine attack began in the morning and in 5 patients in the afternoon.

Table 1 Demographic information of study participants.

	All patients (n=27)	Patients <i>without</i> an attack (n=10)	Patients <i>with</i> an attack (n=17)	Stress sensitive patients (n=9)	Stress insensitive patients (n=8)
Mean age (SD)	40.8 (9.9)	39.1 (10.1)	41.8 (9.9)	41.3 (8.5)	42.3 (11.9)
Ratio of men to women	7 : 20	3 : 7	4 : 13	1 : 8	3 : 8
Ratio MO to MA	20 : 7	7 : 3	13 : 4	8 : 1	5 : 3
Attack frequency per month (SD)	4.4 (2.7)	3.9 (2.1)	4.7 (3.0)	3.7 (2.1)	5.8 (3.6)
PWSQ				58.3 ± 12.5	39.0 ± 9.8*
Cohen PSS				29.4 ± 7.9	16.4 ± 4.2**

MO denotes migraine without aura, MA migraine with aura, PWSQ Penn State Worry Questionnaire and Cohen PSS Cohen Perceived Stress Scale. (*p=0.003 and ** p=0.001).

Baseline characteristics

The demographics of the total study population and the various subgroups are given in table 1. There were nine stress sensitive and eight stress non-sensitive patients. The baseline mean PSWQ and Cohen PSS scores were higher in the stress sensitive patients.

Temporal changes in stress related variables

The temporal profiles of the mean scores for perceived stress, morning cortisol, evening cortisol, heart rate, LF and HF power are shown in Figures 1a-e for the whole study population and in Figures 2a-e for the subgroup of nine stress-sensitive patients compared to eight non stress-sensitive patients. In the total study population, the mean score for perceived stress was 17.8 ± 16.2 on the migraine day, the mean morning cortisol 15.6 ± 9.7 nmol/l, the mean evening cortisol 5.3 ± 2.7 nmol/l and the mean heart rate 79.7 ± 12.1 bpm. Differences between observation days were not significant. The comparison between the stress sensitive with non-sensitive patients revealed in the nine stress sensitive patients an increase in perceived stress in the days prior to an attack (Figure 2a), but no other differences between the two groups.

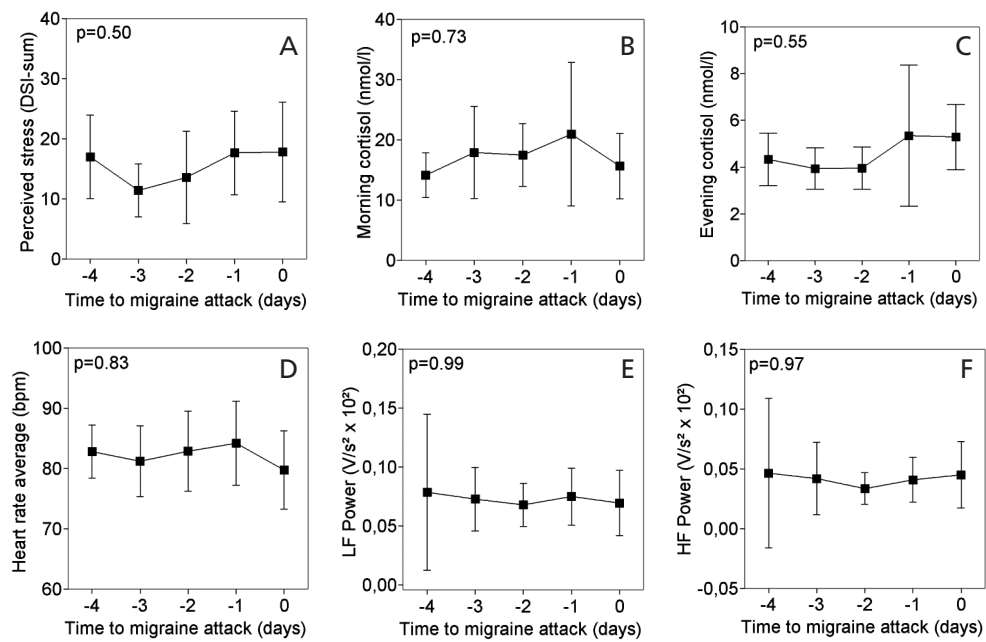


Figure 1A Perceived stress during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 1B** Morning cortisol during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 1c** Evening cortisol during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 1D** Heart rate average during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 1E** LF power during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 1F** HF power during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals.

regel 1
 regel 2
 regel 3
 regel 4
 regel 5
 regel 6
 regel 7
 regel 8
 regel 9
 regel 10
 regel 11
 regel 12
 regel 13
 regel 14
 regel 15
 regel 16
 regel 17
 regel 18
 regel 19
 regel 20
 regel 21
 regel 22
 regel 23
 regel 24
 regel 25
 regel 26
 regel 27
 regel 28
 regel 29
 regel 30
 regel 31
 regel 32
 regel 33
 regel 34
 regel 35
 regel 36
 regel 37
 regel 38
 regel 39

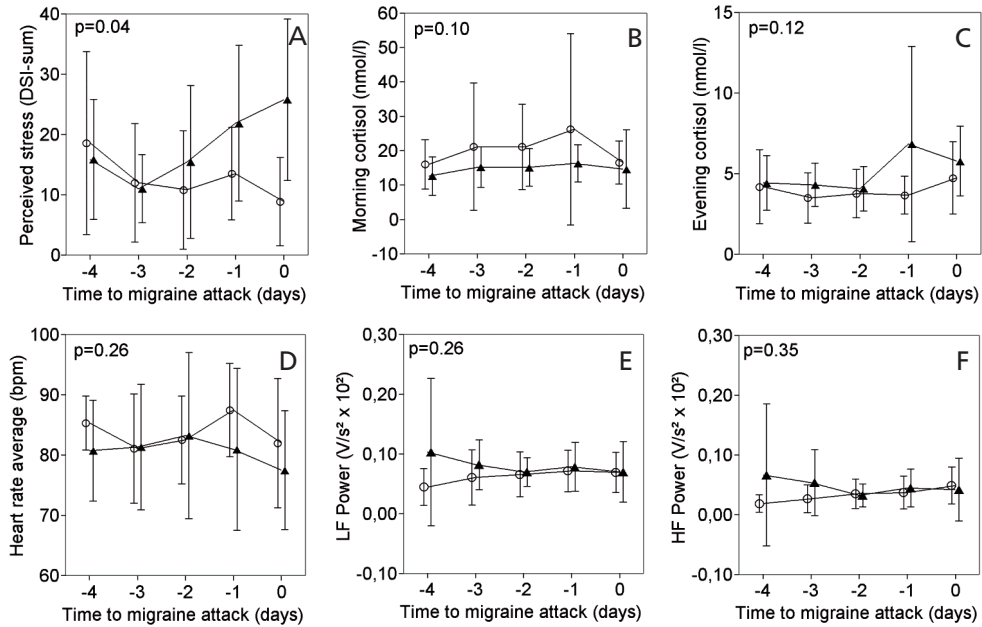


Figure 2A Perceived stress for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 2B** Morning cortisol for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 2C** Evening cortisol for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 2D** Heart rate average for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 2E** Mean LF power for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals. **Figure 2F** Mean HF power for the nine stress sensitive (filled triangles) and eight stress non-sensitive (open circles) patients, who were followed during an attack and for 1-4 days prior to the attack. Error bars represent 95% confidence intervals.

DISCUSSION

In this prospective longitudinal study we failed to find any objective evidence for a temporal relationship between perceived stress, biological indicators for a stress-response, and the onset of migraine attacks. Although stress-sensitive patients indeed reported an increase in perceived stress in the days before an attack, this was not accompanied by objective signs indicating a biological stress response. The present results extend earlier negative findings on the putative relationship between stress and migraine. Autonomic function tests during migraine attacks failed to show changes in heart rate variability, blood pressure reaction¹⁵¹ or transcranial Doppler response in the middle cerebral artery⁴⁴. In contrast, in the inter-ictal phase changes in both sympathetic and parasympatric autonomic function have been described^{95,96,152}. The increase in perceived stress in stress-sensitive patients is in accordance with previous prospective studies in which, however, no biological stress markers were included^{42,43}.

Stressors can be described as physical and psychological events that in potential threaten homeostasis of a living organism¹⁴³. Both acute stressors and stressful daily events have shown to increase cortisol^{153,154} and heart rate¹⁵⁵. Although a profound effect of daily stressful events on migraine seems unlikely, we cannot fully exclude an association between mental stress and migraine. We could only measure 17 migraine patients because of the rather demanding design of the study (daily observations for, in some instances, several weeks because of the unpredictable timing of attacks). Due to the prospective nature of our study the pre-ictal interval varied between study subjects. Twelve out of 17 migraine patient were studied for the full length of 4 days, five patients for a shorter period of time because these five patients experienced their attack within a few days after starting the observation period. Because we did not observe differences for our parameters between day -4 and day -2 we believe that this shorter observation period will not influence our findings. Furthermore, the temporal resolution of our measurements was relatively low. Cortisol was measured only in the morning and evening, and heart rate only in the evening to reduce the effect of physical activity. Theoretically, a reduction in physical activity during evening hours because of the prodromal phase of a migraine attack¹⁵⁶ may have masked an association between changes in heart rate and migraine. Also theoretically, due to the low resolution of measurements this could have resulted in missing changes occurring immediately before the onset of an attack. We feel however that, based on the time course of premonitory symptoms, changes are to be expected to occur 12 to 24 hours prior to the onset of attacks¹³⁴.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____
regel 2 _____
regel 3 _____
regel 4 _____
regel 5 _____
regel 6 _____
regel 7 _____

For our study we excluded pure menstrual migraine. Free salivary cortisol is decreased during the follicular phase of the menstruation period and in oral contraceptive users¹⁵⁷. We did not correct for the temporal relation between menstrual cycle or oral contraceptive use and the occurrence of the migraine attack in the 13 women who were included in this study. Therefore, oral estrogens or the menstrual cycle might have influenced cortisol measurements.

regel 8 _____
regel 9 _____
regel 10 _____
regel 11 _____
regel 12 _____
regel 13 _____
regel 14 _____
regel 15 _____
regel 16 _____
regel 17 _____

Future studies could include continuous measurements including the full 24 hours prior to the onset of attacks, although this will be logistically quite challenging. Although salivary morning cortisol is related to workstress¹⁵⁸, short lasting daily stressors are probably better assessed using high frequent daily measurements¹⁵³. The cortisol response after acute stressors has shown to normalize after 1 to 2 hours¹⁵⁹. Future longitudinal stress studies in migraine could also include epinephrine and norepinephrine as indicators for sympathetic-adrenal-medullary system related changes after mental and physical stressors¹⁶⁰. Both catecholamines can be measured in urine enabling environmental measurements¹⁶¹.

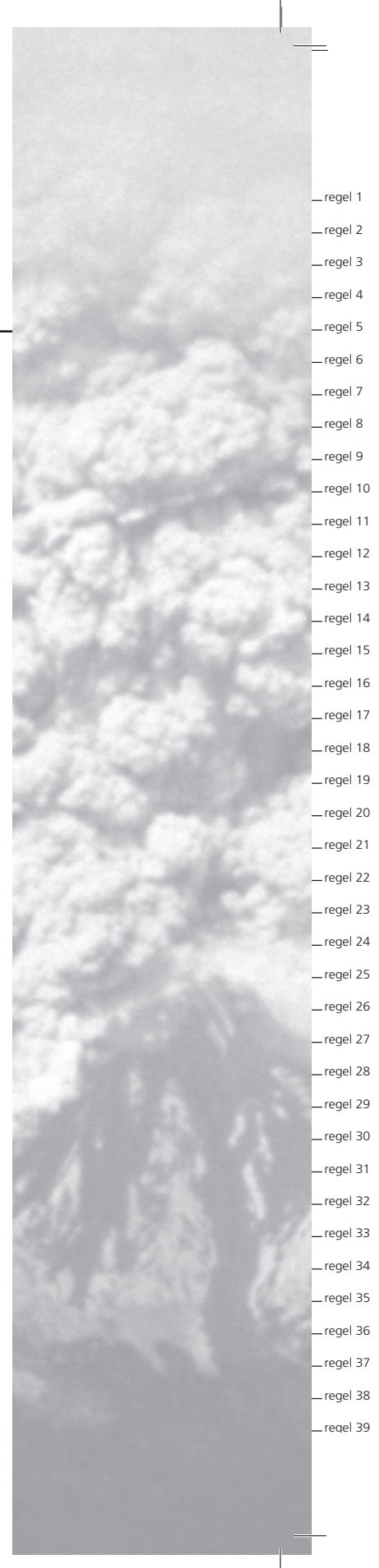
regel 18 _____
regel 19 _____
regel 20 _____
regel 21 _____
regel 22 _____
regel 23 _____
regel 24 _____
regel 25 _____
regel 26 _____
regel 27 _____
regel 28 _____
regel 29 _____
regel 30 _____
regel 31 _____
regel 32 _____
regel 33 _____
regel 34 _____
regel 35 _____
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

In conclusion, we were unable to show objective evidence for a biological stress response before and during migraine attacks. This could reflect a true negative finding or be the result of the discussed study limitations. The reported association between perceived stress and migraine in a sub-population of stress sensitive patients might suggest that these attacks were triggered by mental stress. It could be that in these patients migraine attacks are triggered by mental stress or that events are perceived as stressful due to functional brain changes occurring in the very early phase of a migraine attack.

CHAPTER 3

NORMOBARIC HYPOXIA AND NITROGLYCERIN AS TRIGGER FACTORS FOR MIGRAINE

Cephalalgia 2006;26:816-9



ABSTRACT

Migraine prevalence is increased in high-altitude populations and symptoms of acute mountain sickness mimic migraine symptoms. Here we tested whether normobaric hypoxia may trigger migraine attacks. As positive control we used nitroglycerine (NTG), which has been shown to induce migraine attacks in up to 80% of migraineurs. Sixteen patients (12 females, mean age 28.9 ± 7.2 years) suffering from migraine with ($n=8$) and without aura ($n=8$) underwent 3 different provocations (normobaric hypoxia, NTG and placebo) in a randomized, cross-over, double dummy design. Each provocation was performed on a separate day. The primary outcome measure was the proportion of patients developing a migraine attack according to the criteria of the International Headache Society within 8 hours after provocation onset. Fourteen patients completed all three provocations. Migraine was provoked in 6 (42%) patients by hypoxia, 3 (21%) by NTG and 2 (14%) by placebo. The differences among groups were not significant ($p=0.197$). The median time to attacks was 5 hours. In conclusion, the (remarkably) low response rate to NTG is surprising in view of previous data. Further studies are required to fully establish the potency of hypoxia in triggering migraine attacks.

INTRODUCTION

Migraine is a common neurovascular disorder that affects 15 to 20 % of the population¹¹. Several substances are known to induce migraine attacks in susceptible patients. Nitroglycerin (NTG) is the most frequently studied trigger factor and has been shown to induce migraine attacks in 60 to 80% of migraineurs within 5 to 6 hours^{55,56,78}. Hypoxia may also be a trigger factor for migraine. Firstly, acute exposure to high altitude may induce acute mountain sickness (AMS), which is characterized by headache, insomnia, dizziness, lassitude, fatigue and gastrointestinal symptoms such as anorexia, nausea, or vomiting in an unacclimatized person who has recently reached an altitude above 2500 m⁹⁷. Up to one third of subjects with acute AMS also fulfill the criteria for migraine^{3,98,99}. Secondly, chronic exposure to high altitude is associated with an increased migraine prevalence^{100,101} and thirdly, sumatriptan is an established drug for the acute treatment of migraine⁷⁵, and was also shown to be effective in some studies in AMS^{102,103}. In the present study we tested whether normobaric hypoxia may trigger migraine attacks in migraine patients under experimental conditions. We used NTG as a positive control.

METHODS

Patients

Patients with a history of migraine with (MA) or without (MO) aura, aged 18-65 years, with a baseline attack frequency of 1 to 9 per 3 months in the last six months were recruited from the outpatient clinic, among hospital staff and university students. Exclusion criteria were headache on more than 10 days per month, pregnancy, lactation, psychiatric disorders including substance and drug abuse, neurological diseases other than migraine, and a medical disease that could, according to the judgment of the investigators, interfere with the study. Before each provocation it was made sure that no migraine attack had occurred within the previous 3 days, no pain or migraine medications were taken the previous 24 hours, and that the patient did not suffer from sinusitis or coryza. The study was approved by the local ethical committee.

Experimental Design

Patients were subjected to three different provocations (normobaric hypoxia, NTG and placebo) in a randomized, double-dummy controlled fashion using a cross-over design. The NTG and placebo part were double blind, and the hypoxia part was single blind, because arterial oxygen saturation (SaO₂) had to be monitored continuously.

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

Each of the three provocations was performed on a different day. At the beginning of each provocation, the supine patient obtained a well fitting facial mask, which was connected with a tube for the administration of air with reduced or normal (placebo) oxygen content. Then an antecubital vein was cannulated for the infusion of NTG or saline (placebo). As soon as the patients stated that they became familiar with the facial mask and the attached tube, the provocation was started. An independent physician carried out randomization.

Exposure to normobaric hypoxia: An investigator progressively increased the concentration of nitrogen (N_2) in the inspired air to obtain SaO_2 values of 75 to 80% within 20 minutes. During exposure to normobaric hypoxia, intravenous (IV) saline was administered. The NTG provocation consisted of IV administration of 0.5 microgram / kg body weight NTG within 20 minutes using a free infusion set (Codan, the Netherlands), while the patient was breathing normal air. Placebo provocation: The participants breathed normal air during the whole provocation, whereas only IV saline was administered during the first 20 minutes of the provocation.

Headache Response to the Different Provocations

Migraine symptoms according to the criteria of the International Headache Society (IHS)³ and headache severity on a visual analogue scale (VAS) ranging from 0 to 100 were assessed every 30 minutes. Each provocation was terminated after 5 hours, or earlier, if headache symptoms fulfilled the IHS criteria for migraine, or the experiment was not tolerated by the patient. The presence of headache symptoms was re-assessed 8 hours after the beginning of each provocation, because the time-course of migraine attacks induced by hypoxia might differ from those induced by NTG. After termination of every provocation the patient was asked which provocation they thought they were exposed to.

SaO₂ measurements

SaO_2 was measured using a fingertip pulse oximeter (Datex-Ohmeda, Helsinki, Finland).

Statistical analysis

The primary outcome measure was the migraine response, defined as the proportion of patients developing a migraine attack fulfilling the IHS criteria³ for migraine within 8 hours after the start of the experiment. Differences in response between groups were tested using Friedman's test. Patients who did not complete all provocations were analyzed on a worst-case scenario basis (meaning an attack after placebo and no attack after provocation). Fourteen patients were required to detect a difference in migraine

response of 40% between hypoxia and placebo (alpha 0.05, beta 90%). The secondary outcome measure was the difference in headache response categorized as (1) absent, (2) mild, (3) moderate or severe headache not fulfilling the criteria for migraine or (4) migraine fulfilling the IHS criteria.

RESULTS

A total of 16 patients (12 females, mean age 29 ± 7 years) were included in the study. The mean baseline attack frequency was 1.2 attacks per month (SD 0.76). Fourteen patients completed all three provocations, and two patients completed only two (Table 1).

Table 1 Patient characteristics (demographic and migraine)

Subject	Sex	Age	Migraine (IHS)	Migraine attacks per month	Attack positive provocations	Migraine characteristics of provoked attacks								
						HS	UH	AH	PH	N	V	PT	PN	VAS
1	F	25	MO	0.33	Hypoxia	2	-	-	yes	yes	-	yes	-	59
2	M	26	MA	0.33										
3	M	36	MA	0.33	Placebo	2	-	yes	yes	yes	-	yes	yes	43
4	F	23	MO	3	Hypoxia	3	yes	yes	yes	yes	yes	yes	-	60
5	F	25	MO	1	Hypoxia	3	-	yes	yes	yes	-	yes	-	65
6	M	24	MO	2										
7	F	23	MO	1	Hypoxia	2	-	yes	yes	yes	-	-	-	49
					NTG	2	-	yes	yes	yes	-	-	-	31
8	F	28	MO	1	NTG	2	-	yes	yes	yes	-	-	-	61
9*	F	42	MO	1	NTG	2	yes	yes	yes	yes	-	yes	yes	38
10	F	33	MA	1	Hypoxia	2	yes	yes	yes	yes	-	-	-	70
11*	F	44	MA	2										
12	M	36	MA	2	Hypoxia	2	yes	yes	yes	yes	-	yes	yes	28
					NTG	3	yes	-	yes	yes	yes	yes	yes	29
13	M	23	MO	1										
14	F	29	MA	1										
15	F	22	MA	0.5										
16	F	22	MA	2	Hypoxia	2	yes	yes	yes	-	-	yes	yes	51
					Placebo	3	yes	yes	yes	-	-	yes	yes	61

F denotes female, M male, MA migraine with aura, MO migraine without aura, NTG nitroglycerine, HS headache severity (2=moderate, 3 =severe), UH unilateral headache, AH aggravation of headache during physical activity, PH pulsating headache, N nausea, V vomiting, PT photophobia and PN phonophobia

Out of the 14 patients who underwent all three provocations, six patients (43%; 95% confidence interval (CI) 27% to 69%) developed a MO attack during exposure to normobaric hypoxia, three patients (21%; 95%CI 0% to 42%) after the administration of NTG, and two patients (14%; 95%CI -4% to 32%) after the administration of placebo. The frequency of migraine attacks did not differ among groups ($p= 0.197$). Both patients with incomplete provocations developed a MO attack, one after exposure to normobaric hypoxia and the other after administration of NTG. The inclusion of the two patients who underwent just two provocations did not change the study results ($p=0.150$). The median time to migraine attacks was five hours (4 hours for placebo, 4.5 hours for hypoxia and 6 hours for NTG). Headache responses (Figure 1) did not differ between groups ($p=0.094$). Both in the hypoxia and NTG group there were 4 patients who developed moderate to severe headache, but did not fulfill IHS criteria for migraine (no accompanying symptoms such as nausea, phonophobia or photophobia). The subjects' rating of whether they had been exposed to hypoxia, NTG or placebo was no better than by chance. Four patients guessed all three provocations correct, five guessed all three provocations and four were correct in one provocation (2 placebo and 2 hypoxia). Ratings were missing in one patient.

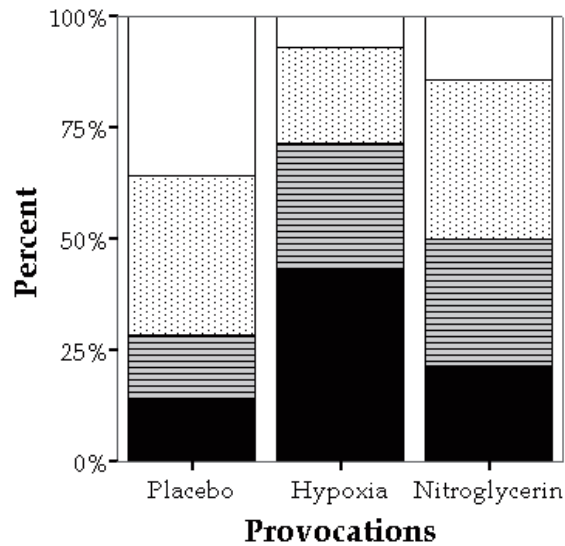


Figure 1 Headache and migraine response to placebo, normobaric hypoxia and nitroglycerin. Black bars represent migraine response, dark gray is moderate or severe headache not fulfilling migraine criteria, light gray is mild headache and white bars is no headache.

DISCUSSION

The first remarkable finding in this study is the low migraine response of 21% after NTG. This is in line with a recent study in English subjects where the migraine response rate after NTG was only 20%¹²³. However in most other studies NTG provoked migraine attacks in 60% to 80% of subjects^{55,78,123,125}. The low response in our study could have been due to either differences in methodology or in study population. Although we administered the same NTG dose and used the same infusion systems (PVC free) as was done in previous studies⁷⁸, the experimental design of our study was entirely different^{55,78}. Due to the double dummy design, the patients had to breathe through a facial mask during the whole duration of all experimental conditions, which was considered rather stressful, but tolerable by most participants. The stress could have prevented the occurrence of migraine attack^{25,26}. Alternatively, our study population could have been less susceptible to NTG. We had 50% of MA patients in our study and such patients may have a lower migraine response to NTG than MO patients^{55,56,162}. Why MA patients would be less susceptible to NTG is not known. A third explanation could be the clinical scoring system. In our study four patients in both the hypoxia and the NTG group had moderate to severe headache but did not fulfill the criteria for migraine.

Normobaric hypoxia provoked a migraine attack in 6 out of 14 patients as compare to only two after placebo and three after NTG. Although this difference between groups was not significant, the relatively high migraine response after hypoxia is remarkably and seems compatible with the results of a large study in mountaineers at high altitude. Of 1213 mountaineers 589 developed headache within 2 to 6 hours after arrival at 4559 m of altitude⁹⁹. In 112 (19%) subjects the symptoms fulfilled the criteria for migraine whereas only 78 (13%) subjects had a history of migraine at sea level. We conclude that the migraine response to NTG was remarkably low in view of previous data, and normobaric hypoxia might be a trigger factor for migraine, but this requires further research.

— regel 1
 — regel 2
 — regel 3
 — regel 4
 — regel 5
 — regel 6
 — regel 7
 — regel 8
 — regel 9
 — regel 10
 — regel 11
 — regel 12
 — regel 13
 — regel 14
 — regel 15
 — regel 16
 — regel 17
 — regel 18
 — regel 19
 — regel 20
 — regel 21
 — regel 22
 — regel 23
 — regel 24
 — regel 25
 — regel 26
 — regel 27
 — regel 28
 — regel 29
 — regel 30
 — regel 31
 — regel 32
 — regel 33
 — regel 34
 — regel 35
 — regel 36
 — regel 37
 — regel 38
 — regel 39

— |

| —

—————

— |

| —

CHAPTER 4

**EXPERIMENTAL HYPOXIA INDUCED
ACUTE MOUNTAIN SICKNESS IS
ASSOCIATED WITH INTRACELLULAR
CEREBRAL OEDEMA.
A 3 TESLA MAGNETIC
RESONANCE IMAGING STUDY**

Journal of Cerebral Blood Flow and Metabolism. 2008;28:198-206



ABSTRACT

Acute mountain sickness is common amongst not acclimatized persons ascending to high-altitude; the underlying mechanism is unknown, but may be related to cerebral edema. Nine healthy male students were studied before and after 6-hours exposure to isobaric hypoxia. Subjects inhaled room air enriched with N₂ to obtain SaO₂ values of 75-80%. Acute mountain sickness was assessed with the environmental symptom questionnaire, and cerebral edema with 3 Tesla magnetic resonance imaging in 18 regions of interest in the cerebral white matter. The main outcome measures were development of intra- and extracellular cerebral white matter edema assessed by visual inspection and quantitative analysis of apparent diffusion coefficients, derived from diffusion-weighted imaging, and B0 signal intensities, derived from T2-weighted imaging. Seven of nine subjects developed acute mountain sickness. Mean apparent diffusion coefficient increased 2.12% (baseline, 0.80 ± 0.09; 6-hours hypoxia, 0.81 ± 0.09; p=0.034), and mean B0 signal intensity increased 4.56% (baseline, 432.1 ± 98.2; 6-hours hypoxia, 450.7 ± 102.5; p<0.001). Visual inspection of magnetic resonance images failed to reveal cerebral edema. Cerebral acute mountain sickness scores showed a negative correlation with relative changes of apparent diffusion coefficients (r=-0.83, p=0.006); there was no correlation with relative changes of B0 signal intensities. In conclusion, isobaric hypoxia is associated with mild extracellular (vasogenic) cerebral edema irrespective of the presence of acute mountain sickness in most subjects, and severe acute mountain sickness with additional mild intracellular (cytotoxic) cerebral edema.

INTRODUCTION

Unacclimatized subjects, who rapidly ascent to high-altitude, may develop acute mountain sickness (AMS) that is characterized by headache, anorexia, nausea, vomiting, insomnia and dizziness^{97,163-165}. The underlying mechanism for AMS is unclear. Intracellular (cytotoxic) cerebral edema, extracellular (vasogenic) cerebral edema, and increased cerebral blood volume have all been implicated, but without convincing scientific evidence^{97,164,166}. Some magnetic resonance imaging (MRI) studies, using 1.5 Tesla machines, found that exposure to moderate hypo- or isobaric hypoxia, corresponding to altitudes of 4500 m, increased brain volume by 0.5-2.8%^{167,168} and decreased cerebrospinal fluid volume in the lateral and third ventricles by 10.3%¹⁶⁹. Results with respect to the presence and type of cerebral edema were, however, conflicting¹⁶⁷⁻¹⁷⁰. This could have been related to the relatively low resolution of 1.5 Tesla MRI and because the MR images were only visually evaluated and not analyzed with more sensitive quantitative methods. In the present study, we used 3 Tesla MRI to investigate whether experimental hypoxia-induced AMS is associated with intra- and/or extracellular cerebral edema. We compared diffusion-weighted (DWI) and T2-weighted (T2WI) MR images obtained before and after 6-hours exposure to isobaric hypoxia by visual inspection and quantitative analysis of apparent diffusion coefficients (ADCs) derived from DWI, and B0 signal intensities derived from T2WI.

MATERIALS AND METHODS

Subjects

Nine healthy male volunteers (mean age 26.4 ± 3.5 years) were recruited from students of the University of Zurich, Switzerland. Exclusion criteria were: altitude exposure (>1500 m) in the previous 3 months, a history of smoking, substance and drug abuse, or of lung, cardiac, neurological or psychiatric disease, and contraindication to undergo MRI (e.g., pacemaker).

The local medical ethical committee approved the study protocol, and written informed consent was obtained from all subjects.

Study design

The study subjects were in supine position and were fitted a facial mask which was connected with a tube of 3 m length. The total flow of fresh gas varied between 9 to 12 liters per minute, because 6 liters of compressed air per minute were mixed with

— regel 1

— regel 2

— regel 3

— regel 4

— regel 5

— regel 6

— regel 7

— regel 8

— regel 9

— regel 10

— regel 11

— regel 12

— regel 13

— regel 14

— regel 15

— regel 16

— regel 17

— regel 18

— regel 19

— regel 20

— regel 21

— regel 22

— regel 23

— regel 24

— regel 25

— regel 26

— regel 27

— regel 28

— regel 29

— regel 30

— regel 31

— regel 32

— regel 33

— regel 34

— regel 35

— regel 36

— regel 37

— regel 38

— regel 39

3 to 6 liters of N₂ per minute. Thus, the stimulus was assumed to cause poikilocapnic hypoxia. Baseline examinations were done when the subjects were familiarized with the facial mask and the attached tube, and included assessment of the cerebral symptoms of AMS (AMS-C) by completion of the environmental symptom questionnaire (ESQ)¹⁷¹, monitoring of arterial O₂ saturation (SaO₂), the measurement of the blood pressure, and baseline MRI. The supine subjects were then transported to a room adjacent to the MR suite, and the tube was connected with a gas container. Here, one investigator gradually increased the concentration of N₂ in the inspired air, over a period of 20 minutes, to obtain SaO₂ values of 75 to 80%. This corresponds to an altitude of about 4500 m. The SaO₂ values were measured using a fingertip pulse oximeter (Datex-Ohmeda, Helsinki, Finland). End-tidal pCO₂ was not determined for technical reasons. The SaO₂ was kept stable during the following 6 hours and the second MRI study by adjusting the mixture of inspired gas. After 6 hours of hypoxia, the subjects were transported in the supine position to the MR suite for the second MRI study. Symptoms of AMS and blood pressure were re-assessed every hour, and finally during the second MRI study.

MRI studies

MRI studies were performed using a 3 T Philips Intera whole body system (Philips Medical Systems, Best, The Netherlands). Identical protocols and volume positioning were used at baseline and during hypoxia. The DWI data were based on a spin-echo single excitation echo-planar imaging protocol. Whole brain scans with an in-plane resolution of 1.6 x 1.6 mm² (14 contiguous slices, slice thickness = 4 mm, matrix = 128², echo time = 79 ms, relaxation time = 3987 ms) were carried out along three orthogonal diffusion directions with a diffusion weighting of $b = 1000 \text{ s/mm}^2$, of $b=0$ ¹⁷² and of B0 images (T2* weighted images from the same sequence, with no applied diffusion gradient). An isotropic diffusion-weighted image (Figure 3A) was calculated as the geometric mean of three orthogonal diffusion-weighted images. Additionally, for each slice a T2WI with minimal diffusion weighting ($b < 20 \text{ s/mm}^2$) was acquired. The duration for the imaging procedure including the diffusion and the T2 protocol was 4 min and 6 s.

The MRI data were stored and independently analyzed after completion of the study by investigators who were not aware of the cerebral AMS (AMS-C) scores. Two physicists (TJ and UD) and neurologists (PSS and RWB), blinded to the AMS scores, looked for the presence of cerebral edema by comparing the second DWI and T2WI scans of each subject with the corresponding baseline scans (Figure 3). Another neurologist (ACN), also blinded to the AMS scores, measured the ADC values and B0 signal intensities in 22 regions of interest (ROI) as the average value of all pixels in the respective ROI.

The ROIs were circular and located on four consecutive slices (Figure 1). Slice A and B

were placed above, and the next two slices at the level of the cella media of the lateral ventricles. Eighteen of 22 ROIs were located in the cerebral white matter (nine in each hemisphere), and the other four in the cerebrospinal fluid (CSF) of both lateral ventricles. Cerebral white matter ROI were located in the anterior part in eight cases (beside the forceps minor; two measurements per slice), the middle part for two ROIs (beside the lateral ventricles; two measurements on the lowest slice), and the posterior part for eight ROIs (next to the forceps major, which corresponds to the lateral part of the corpus callosum fibers; two measurements per slice). Each white matter ROI consisted of 88 pixels, and each CSF ROI of 16 pixels. CSF ROIs contained fewer pixels than ROIs in the white matter, because not all ventricles were wide enough to accommodate larger circles. The use of smaller reference ROI in the CSF is appropriate, since the MRI signal is higher in the CSF than the white matter and thus provides a better signal-to-noise ratio. MRI signal intensities are arbitrary units with different absolute values at baseline and 6-hours sessions. Therefore, ROIs placed in the CSF were used to correct for intersession differences, because the CSF signal levels are assumed to remain unchanged during hypoxia. All ADC values and B0 signal intensities measured after 6-hours were thus corrected to achieve the same values in CSF as at baseline according to the proportion: $ROI_{6\text{-hours corrected}} = ROI_{6\text{-hours}} \times CSF_{\text{baseline}} / CSF_{6\text{-hours}}$, where CSF is the mean value of all CSF ROIs.

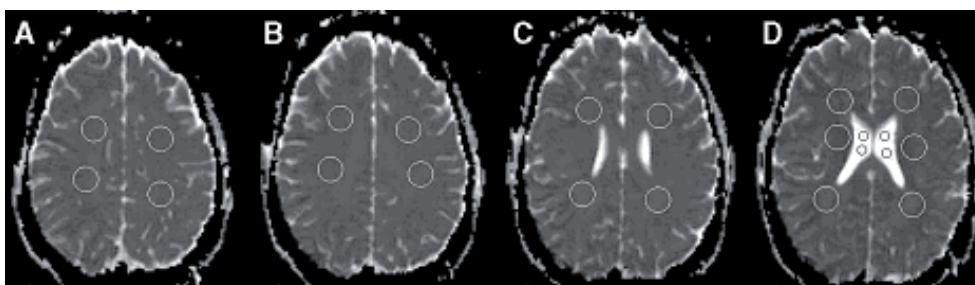


Figure 1 Axial T2-weighted MR images show the 22 regions of interest. Slice A placed above slice B, and slices C and D placed at the level of the cella media of the lateral ventricles.

Interpretation of ADC and B0 changes on MRI

Increase of both ADC and B0 values are indicative of extracellular (vasogenic) edema, whilst an increase of B0 values in combination with a decrease of ADC values is indicative for the development of intracellular (cytotoxic) edema ¹⁷³⁻¹⁷⁷.

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

Assessment of AMS

The ESQ was translated to German and used as described previously¹⁷⁸. Subjects were considered to suffer from AMS when the AMS-C score was ≥ 0.70 ¹⁷¹. The AMS-C score ranges from 0-5 and is based on eleven neurological symptoms.

Statistical analysis

Statistical analysis was performed using SPSS 12.0 (SPSS, Chicago, Illinois). Mean ADC (primary outcome of the study) and B0 values obtained at baseline and after 6 hours were compared using a general linear model for repeated measurements including ROI location as covariate (total white matter ROI, n=162; anterior white matter ROI, n=72; middle white matter ROI, n=18; posterior white matter ROI, n=72). Associations between AMS-C scores and relative changes of ADCs and B0 signal intensities were assessed using the non-parametric Spearman correlation coefficients. P <0.05 was considered significant.

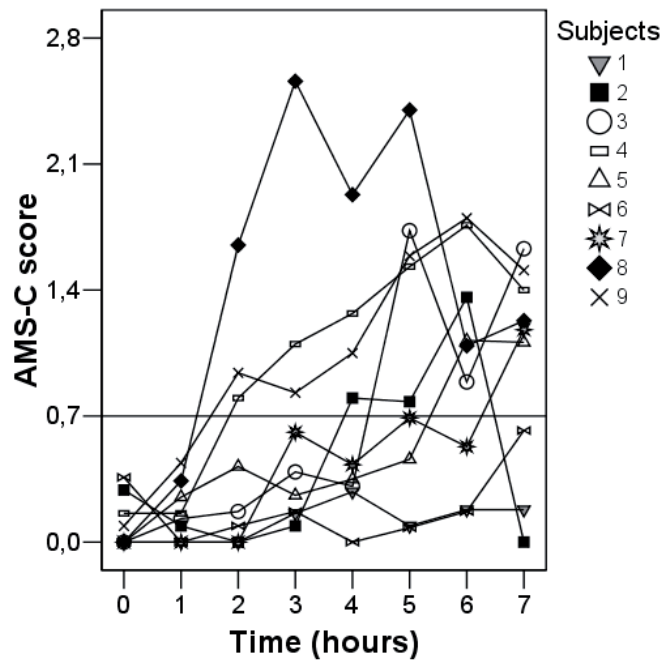


Figure 2 Cerebral acute mountain sickness (AMS-C) scores of all subjects at baseline and during hypoxia.

RESULTS

All nine subjects completed the study. The data set was complete and was evaluated for all 9 subjects. Seven out of nine subjects (subjects 2-6, 8, 9) developed AMS during hypoxia. Six of the nine subjects had AMS during the second MRI scan (Figure 2). Baseline blood pressure did not differ between subjects with and without AMS. The mean AMS score of all subjects was higher at the time of MRI scanning than at baseline (Table 1). Systolic (baseline, 113 ± 9 mm Hg; during second MRI study, 115 ± 11 mm Hg; $p=0.49$) and diastolic (baseline, 72 ± 6 mm Hg; during second MRI study, 74 ± 4 mm Hg; $p=0.71$) blood pressure did not change during the study (values are means \pm standard deviation).

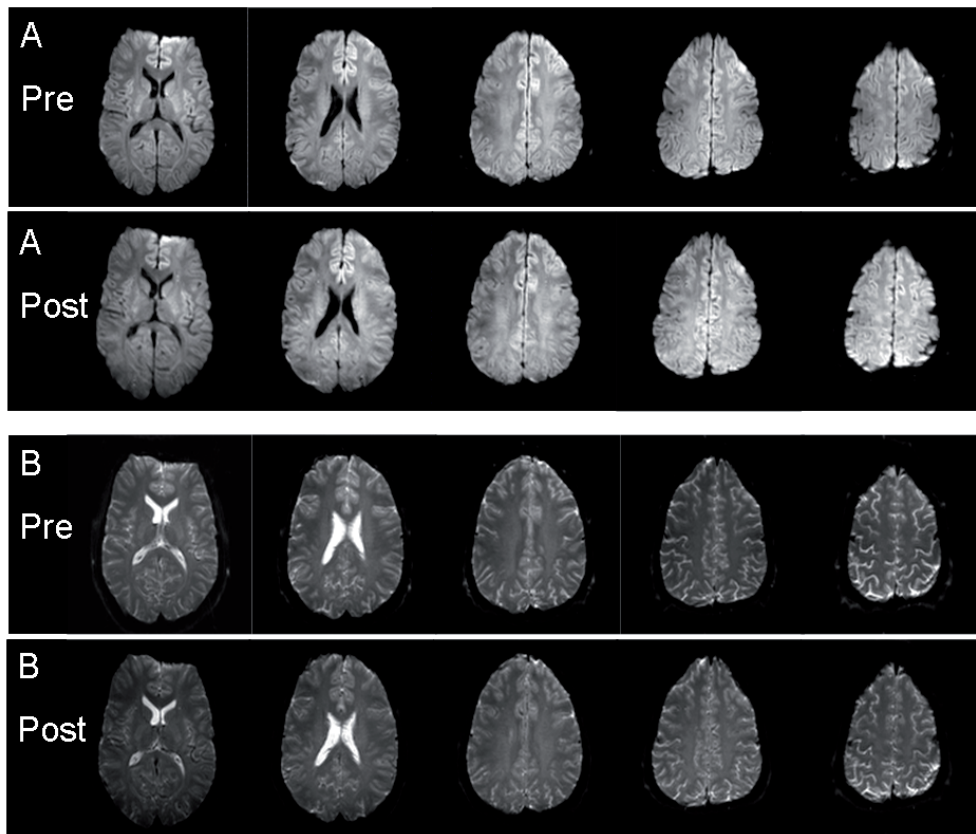


Figure 3 Axial diffusion-weighted (A) and T2-weighted (B) MR images obtained at baseline (upper rows) and after 6-hours exposure to hypoxia (lower rows) from subject 4 who suffered from severe acute mountain sickness. There is no evidence for the development of cerebral edema at visual inspection.

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

Table 1 Apparent Diffusion Coefficient Values at 3 Tesla MR Imaging in the Cerebral White Matter and Cerebrospinal Fluid, and Cerebral Acute Mountain Sickness Scores in Nine Healthy Subjects at Baseline and 6-Hours Exposure to Isobaric Hypoxia*

ADC		Cerebral White Matter		CSF†		AMS-C score					
		Anterior		Middle		Posterior		All		Score	
		Baseline	% Change at 6-h hypoxia†	Baseline	% Change at 6-h hypoxia†	Baseline	% Change at 6-h hypoxia†	Baseline	% Change at 6-h hypoxia†	Baseline	After 6-h hypoxia
Subject 1	0.70	0.70	7.72	0.66	-0.79	0.70	6.98	0.69	6.45	0.00	0.18
Subject 2	0.72	0.72	-5.11	0.71	-4.26	0.76	-3.40	0.74	-4.25	0.29	1.36
Subject 3	0.78	0.78	3.47	0.72	5.45	0.81	4.90	0.79	4.32	0.00	0.89
Subject 4	0.95	0.95	-7.27	0.92	-3.98	0.96	-6.53	0.95	-6.58	0.16	1.76
Subject 5	0.78	0.78	2.57	0.70	10.93	0.78	3.57	0.77	3.94	0.00	1.12
Subject 6	0.74	0.74	9.27	0.69	5.99	0.76	10.84	0.74	9.60	0.36	0.17
Subject 7	0.92	0.92	3.12	0.91	4.01	0.96	3.40	0.94	3.34	0.00	0.53
Subject 8	0.80	0.80	1.78	0.78	3.17	0.84	5.02	3.29	3.37	0.00	1.09
Subject 9	0.74	0.74	-0.20	0.70	0.06	0.75	-2.26	0.74	-1.09	0.09	1.80
Subjects 1-9 mean	0.79	0.79	1.71	0.75	2.29	0.81	2.50	0.80	2.12	0.10	0.99
SEM	0.03	0.03	0.90	0.03	1.20	0.03	0.88	0.09	0.57	0.05	0.20
Difference baseline vs 6 h at hypoxia								p=0.034			p=<0.01

h denotes hours, and SEM standard error of the mean. * The concentration of N₂ in the inspired air was adjusted to obtain arterial SO₂ values of 75 to 80%. † The apparent diffusion coefficient values obtained before exposure to isobaric hypoxia were defined as 100% † The values for the relative change of the apparent diffusion coefficient measured in the cerebrospinal fluid after 6-hours of isobaric hypoxia were zero, because they were corrected to achieve the same values as at baseline (for details see text).

MRI study

Visual inspection showed no evidence for cerebral edema (Figure 3).

Mean ADCs were increased by 2.12% ($p=0.034$, Table 1), and mean B0 values were increased by 4.56% ($p<0.01$, Table 2) after 6-hours exposure to hypoxia. The ADCs increased in 6 subjects (Figure 4), and B0 values in 8 subjects (Figure 5). ADCs ($p=0.32$) and B0 values ($p=0.06$) did not differ between the 3 white matter ROIs.

As shown in Figure 6, the AMS-C scores measured after 6-hours exposure to hypoxia showed a negative correlation with the relative change of ADC values (Spearman correlation coefficient -0.83 , $p = 0.006$). There was no association between the AMS-C scores measured after 6-hours exposure to hypoxia and the relative change of B0 values (Spearman correlation coefficient -0.22 , $p = 0.576$; Figure 7).

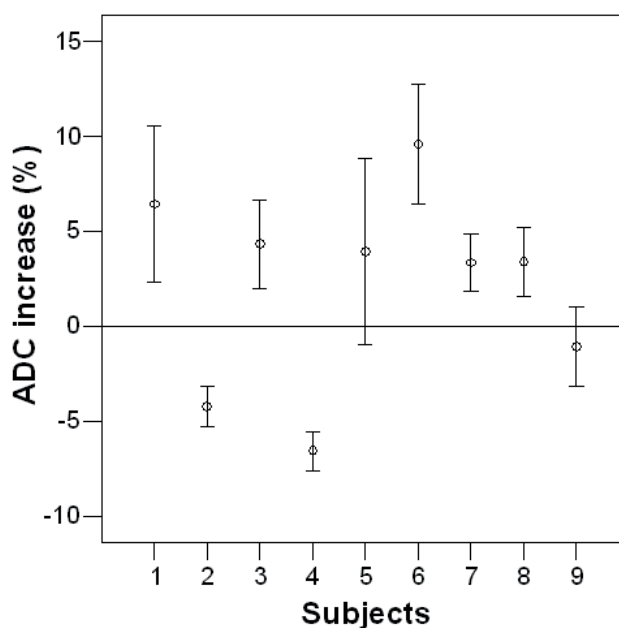


Figure 4 Relative changes of the apparent diffusion coefficient (ADC) after 6 hours of hypoxia compared to baseline in all nine subjects.

regel 1
 regel 2
 regel 3
 regel 4
 regel 5
 regel 6
 regel 7
 regel 8
 regel 9
 regel 10
 regel 11
 regel 12
 regel 13
 regel 14
 regel 15
 regel 16
 regel 17
 regel 18
 regel 19
 regel 20
 regel 21
 regel 22
 regel 23
 regel 24
 regel 25
 regel 26
 regel 27
 regel 28
 regel 29
 regel 30
 regel 31
 regel 32
 regel 33
 regel 34
 regel 35
 regel 36
 regel 37
 regel 38
 regel 39

Table 2 B0 Values at 3 Tesla MR Imaging in the Cerebral White Matter and Cerebrospinal Fluid, and Cerebral Acute Mountain Sickness Scores in Nine Healthy Subjects at Baseline and 6-Hours Exposure to Isobaric Hypoxia*

	AMS-C score										
	CSF [†]										
	Anterior			Middle			Posterior			All	
B0 value	Baseline	% Change at 6-h hypoxia [†]	Baseline	% Change at 6-h hypoxia [†]	Baseline	% Change at 6-h hypoxia [†]	Baseline	% Change at 6-h hypoxia [†]	Baseline	% Change at 6-h hypoxia [†]	Score
Subject 1	186.13	8.27	224.00	8.11	216.38	2.13	203.78	5.52	879.00	0.00	0.18
Subject 2	354.25	0.42	389.00	6.74	370.75	11.36	365.44	5.98	1353.75	0.29	1.36
Subject 3	452.38	10.86	489.50	12.37	441.75	11.10	451.78	11.13	1630.75	0.00	0.89
Subject 4	475.75	-4.97	488.00	1.16	490.50	0.93	483.67	-1.67	1544.00	0.16	1.76
Subject 5	489.63	5.23	569.00	6.54	563.88	4.08	531.44	4.87	2019.50	0.00	1.12
Subject 6	411.13	1.61	445.00	3.89	430.13	7.77	423.33	4.60	1477.75	0.36	0.17
Subject 7	469.25	1.92	502.50	3.50	508.63	0.80	490.44	1.60	1580.75	0.00	0.53
Subject 8	478.38	4.37	498.50	1.60	464.50	11.07	474.44	7.04	1724.00	0.00	1.09
Subject 9	439.50	6.03	504.00	7.09	479.13	-3.46	464.28	1.93	1373.75	0.09	1.80
Subjects 1-9 mean	417.38	3.75	456.61	5.67	440.63	5.09	432.07	4.56	1509.25	0.10	0.99
SEM	32.11	1.55	33.25	1.18	33.26	1.81	32.45	1.22	103.30	0.05	0.20
Difference baseline vs 6 h at hypoxia											
											p=<0.01

h denotes hours, and SEM standard error of the mean. * The concentration of N₂ in the inspired air was adjusted to obtain arterial SO₂ values of 75 to 80%. [†] The apparent diffusion coefficient values obtained before exposure to isobaric hypoxia were defined as 100% [†] The values for the relative change of the apparent diffusion coefficient measured in the cerebrospinal fluid after 6-hours of isobaric hypoxia were zero, because they were corrected to achieve the same values as at baseline (for details see text).

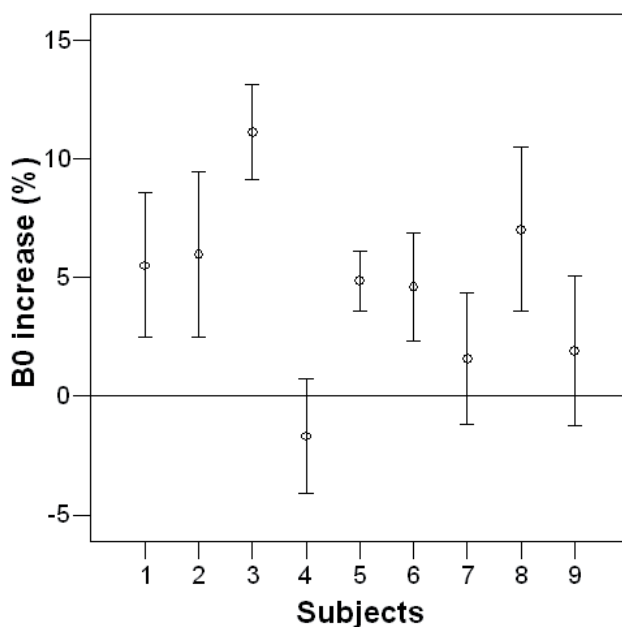


Figure 5 Relative changes of the B0 values after 6 hours of hypoxia compared to baseline in all nine subjects.

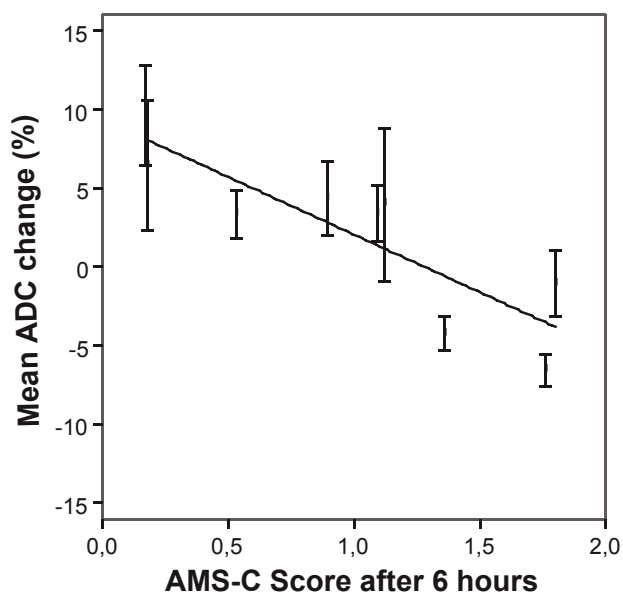


Figure 6 The mean value of two cerebral acute mountain sickness (AMS-C) scores measured immediately before and after MR imaging performed at hypoxia showed a weak but significant negative correlation with the relative change of the apparent diffusion coefficient (ADC).

diffusion coefficient measured in the cerebellum immediately after 6 hours of isobaric hypoxia were zero, because they were corrected to achieve the same values as at baseline (for details see text).

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

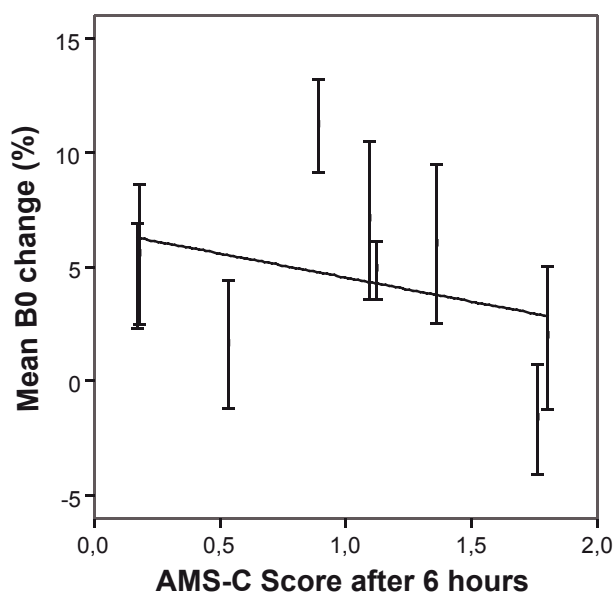


Figure 7 The mean value of two cerebral acute mountain sickness (AMS-C) scores measured immediately before and after MR imaging performed at hypoxia showed no correlation with the relative change of the B0 signal intensity values.

DISCUSSION

In the present 3 Tesla MRI study, we found that experimental isobaric hypoxia for six hours: 1) caused AMS in seven (77%) of nine healthy volunteers; 2) produced a mild extracellular (vasogenic) cerebral edema, irrespective of the presence of AMS, which was identified by a small increase of both ADCs and B0 values, whereas visual inspection of the MRI data failed to detect any differences; and 3) that the AMS-C scores were negatively correlated with the ADC values. The prevalence of AMS in this series is in accordance with the results of two previous studies exposing 31 subjects to isobaric hypoxia corresponding to altitudes of 4500-4564 m during 9-16 hours ^{167,179}. In the study of ¹⁷⁹, 6 (67%) of 9 subjects developed AMS, and in the study of ¹⁶⁷ 11 (50%) of 22 subjects were affected by this altitude illness.

The fact that the mild extracellular (vasogenic) cerebral edema was just detected by quantitative but not visual analysis of the MRI data is in accordance with the results obtained in three 1.5 T MRI investigations ¹⁶⁷⁻¹⁶⁹. These studies found no cerebral edema at visual inspection of T2WI and DWI in 41 subjects with mild to moderate AMS who were exposed to hypobaric or isobaric hypoxia corresponding to altitudes of 4500-

4572 m for 10 to 32 hours¹⁶⁷⁻¹⁶⁹. Furthermore, one of these 3 studies found a mild extracellular cerebral edema with increased B0 values and a trend for decreased ADCs¹⁶⁷. The higher resolution and signal-to-noise ratio of 3 Tesla MRI used in the present study makes it more sensitive for detecting cerebral abnormalities¹⁸⁰ than 1.5 Tesla MRI employed in previous AMS investigations¹⁶⁷⁻¹⁶⁹. Despite this, visual inspection even of 3 Tesla MRI brain images was still not sensitive enough to detect the cerebral edema associated with AMS. This is in sharp contrast to high altitude cerebral edema (HACE). Here visual inspection of proton density- and T2-weighted MRI brain images revealed extracellular edema of the white cerebral matter at a mean of 58 hours (range, 16 to 132 hours) after the onset of HACE symptoms¹⁰⁴.

There was no association between AMS-C scores and B0 signal intensities in this series, which confirms the results of a previous study¹⁶⁷. In this respect it is important to note that, in reality, the degree of extracellular brain edema might have been higher. The B0 signal intensity increase due to cerebral edema may have been neutralized by the blood-oxygenation-level-dependent (BOLD) effect of hypoxia^{181,182}. This effect is related to the intravascular concentration of deoxyhemoglobin, which lowers signal intensity of B0 images by increasing magnetic susceptibility¹⁸³⁻¹⁸⁶. As hypoxia will increase the intravascular concentration of deoxyhemoglobin, it will also lower the intensity of the B0 signal and thus the level of perceived cerebral edema. Therefore, the BOLD effect might have prevented the detection of an association between the B0 values and the AMS-C scores. Consequently, it cannot be completely excluded that the mild extracellular cerebral edema is in part responsible for AMS, e.g. by stimulating pain sensitive fibers in the meninges, the meningeal and pial vessels¹⁸⁷. A potential role of vasogenic cerebral edema is underscored by two observations: Symptoms and signs of AMS as well as abnormal B0 values and ADCs occurring during exposure to isobaric hypoxia disappeared after the subjects were re-exposed to normoxia¹⁶⁷, and corticosteroids, which reduce extracellular cerebral edema, are an established therapy of AMS¹⁸⁸.

The third and most important result of the present study is based on the observation that subjects with the most severe AMS symptoms showed the lowest ADC values. Being in accordance with findings reported by¹⁶⁷, this would suggest that severe AMS is associated with intracellular (cytotoxic) edema of the cerebral white matter, on top of hypoxia-driven extracellular (vasogenic) edema. The cytotoxic edema may have been caused by a decreased activity and/or expression of the Na⁺, K⁺-ATPase in the cerebral white matter¹⁸⁹⁻¹⁹¹. A reduction of ATPase activity is associated with reduced levels of tissue ATP and a shift from aerobic to anaerobic glycolysis, and lactate is built up causing acidosis^{189,191,192}. Using declining intracellular pH (pHi) as an indicator of increased intracellular lactate production, ³¹P MR spectroscopy (MRS) studies observed a pHi

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

decline at arterial PO_2 (PaO_2) values of 30-45 mm Hg^{189,191,193}. We observed PaO_2 values of 40-45 mm Hg in subjects exposed to an altitude of 4559 m^{178,194}. Consequently, the subjects investigated in this series were exposed to levels of hypoxia, which might lead to PaO_2 values causing anaerobic glycolysis and reducing the cerebral metabolic rate of oxygen ($CMRO_2$), although no study has shown a decrease of $CMRO_2$ in humans exposed to high altitude^{195,196}. However, the study of¹⁹⁶ was performed at a lower altitude of 3800 m, and the investigation of¹⁹⁵ in subjects who were chronically exposed to high altitude. More important is that a recent study comparing MRI with positron emission tomography (PET) findings in patients with acute ischemic stroke has shown that ADCs are not reliable predictors of reduced $CMRO_2$ at the levels of hypoxia applied in our subjects¹⁹⁷. These PET findings question the assumption that more severe forms of AMS are associated with intracellular (cytotoxic) edema. Further studies assessing also the $CMRO_2$ in the cerebral white matter are needed to answer this question.

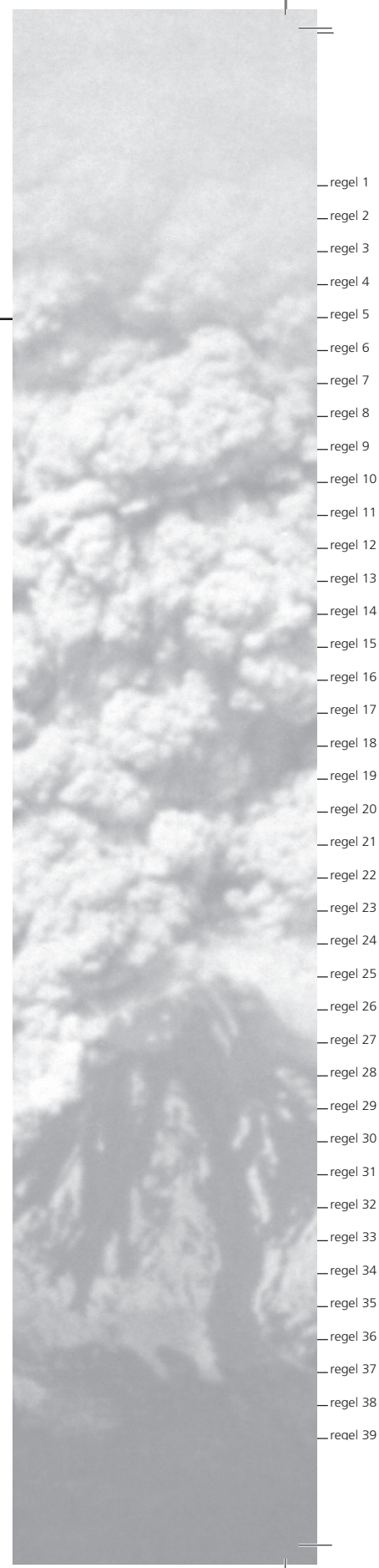
The study is limited by the low number of included subjects. Furthermore, the present findings may not be applicable to hypobaric hypoxia, because the severity of AMS has been shown to be increased during simulated altitude compared with isobaric hypoxia¹⁷⁹.

We conclude that experimental isobaric hypoxia is associated with mild extracellular (vasogenic) cerebral edema irrespective of the presence of AMS in the majority of subjects, and severe AMS with additional mild intracellular (cytotoxic) cerebral edema.

CHAPTER 5

MAGNETIC RESONANCE ANGIOGRAPHY OF THE HUMAN MIDDLE MENINGEAL ARTERY: IMPLICATIONS FOR MIGRAINE

Journal of Magnetic Resonance Imaging. 2006; 24: 918-21



ABSTRACT

Purpose

To describe a novel non-invasive method to study MMA diameter changes *in vivo* in humans. Dilatation of the middle meningeal artery (MMA) has been implicated in the pathophysiology of migraine headache but without direct evidence in humans.

Materials and methods

The diameter of the MMA (extracranial part) was measured in 19 healthy volunteers before and after administration of a vasodilator (nitroglycerin 1.2mg sublingual) known to provoke headache. We used magnetic resonance angiography (MRA) in combination with a 47mm microscopy coil and a semi-automatic contour detection program.

Results

The diameter of the MMA was 1.5 ± 0.26 mm (mean \pm SD) before and 1.79 ± 0.30 mm after nitroglycerin administration. This increase was 20.1% (95% CI 12.9 to 27.3; $p < 0.001$). The mean increase in subjects who developed headache ($n=11$) was 0.34 ± 0.19 mm as compared to $0.22 \text{ mm} \pm 0.20$ mm in the 8 subjects who did not (95% CI for difference: -0.07 to 0.31; $p=0.188$).

Conclusion

MRA in combination with a 47mm microscopy coil is a novel, non-invasive method to measure diameter changes of human meningeal vessels with potential applications in migraine and other neurovascular research.

INTRODUCTION

Migraine is a common and disabling, multifactorial neurovascular headache syndrome^{8,11}. The middle meningeal artery (MMA) has been implicated in the pathogenesis of migraine headache. The dura mater is a pain sensitive structure and mechanical stimulation of the MMA causes pounding migraine-like headache¹⁹⁸.

Sumatriptan is effective in the acute treatment of migraine¹⁹⁹ and may constrict the MMA as demonstrated by selective angiography²⁰⁰. Direct evidence, in humans, for the role of the MMA in migraine headache is, however, lacking. A major reason is that, due to its small diameter (less than 1.86 ± 0.60 mm)²⁰¹, there were no reliable non-invasive methods to measure the MMA *in vivo*. Here we present a Magnetic Resonance Angiography (MRA) based method to non-invasively monitor diameter changes of the MMA. To provoke dilatation of the MMA we used nitroglycerin which is a strong vasodilator and is known to cause migraine headache in up to 60% of migraineurs. In spite of the advantages of contrast enhanced MRA (CE-MRA), we used a non CE-MRA acquisition technique because of medical ethical concerns: in a CE-MRA protocol gadolinium contrast should be delivered twice in relatively short time (less than 30 minutes) with administration of nitroglycerin in between.

METHODS

Subjects

We recruited 22 healthy volunteers (age 18 - 65 years) by public announcement. Exclusion criteria were (A) a history of vascular disease, migraine or any other primary headache syndrome, (B) headache on more than 6 days per month, (C) current use of vasoactive medication, (D) use of more than 3 units of caffeine per day and (E) active smoking. The study was approved by the local ethical committee.

Experimental design

Subjects were asked to refrain from drinking alcohol 24 hours and caffeine containing beverages 12 hours prior to the experiments. MRI scans were performed before and shortly after sublingual administration of NTG 1.2 milligram²⁰². Subjects remained in the MRI scanner and kept their position between scan 1 and 2 to ensure a constant localisation of the measurement. Blood pressure and heart rate were monitored during the experiment. Migraine symptoms were assessed before and after the experiments using the criteria of the IHS³.

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

Magnetic resonance angiography

MRA of the MMA was performed on a 1.5-T system (Philips Medical Systems, Best, the Netherlands). Subjects were positioned using flexible head restraints to minimise the influence of subject movement. Once the MMA was localised using the standard head coil, a small surface coil with a diameter of 47 mm was positioned over the MMA-region and high-resolution MRA images of the MMA were collected. In general the centre of the surface coil was positioned over the Temporo-Mandibular Joint. At this location the MMA is at a depth of around 3 to 4 cm from the skin. The MRA imaging protocol consisted of a sequential 2D acquisition time-of-flight T1-weighted Fast Field Echo MRA sequence with the following imaging parameters: repetition time/echo time, 28 ms/8.7 ms; flip angle, 20°; field of view, 100x100 mm; matrix size, 256 x 256; reconstruction matrix, 256x256; 0.39 x 0.39-mm pixel resolution (0.15-mm² pixel area); number of excitations: 2; slice thickness, 2.0 mm; slices overlapped 1.0 mm; number of slices, 40; total acquisition time 4 min 26 sec. In this scan protocol we applied relatively thick overlapping slices. This is because the image post processing tool makes use of a single 2D slice in which should contain the entire MMA length of interest. Since we expect a MMA diameter of about 1.4-1.5mm, the current scan protocol (2mm slice thickness-1mm overlap) avoids potential partial volume effects.

Image post processing and diameter calculations

All MRA images were transferred to a remote workstation for quantitative analysis using the MR Analytical System (MRI-MASS)²⁰³. The measurement procedure consisted of the manual identification of the borders of the vessel segment to be analyzed. The exact vessel boundaries were detected using an automated contour detection technique based on dynamic programming. The diameter of the vessel segment was automatically derived from the detected vessel contours. MMA-ex was measured in a segment of 7 mm (ranging between 6.5 and 7.5mm), approximately 10 mm from the origo of the Maxillary artery. The 18 diameter measurements that were obtained within the segment (one at every pixel position) were averaged to obtain a mean diameter for the segment. By obtaining multiple measurements the measurement precision could be improved to $0.39 / \sqrt{18} = 0.09$ mm. Reliability of the semi-automatic measurements was also assessed by a second independent observer and agreement between observers was measured by the intra class correlation.

Statistical analysis

The diameter of the MMA before and after nitroglycerin administration was compared with a paired t-test. Differences between subjects with and without headache were

compared with an unpaired t-test. A p value of <0.05 was considered statistically significant.

Sample size calculations

The minimally expected increase of the MMA during migraine headache is unknown. Friberg estimated a mean 20% increase of the diameter of the middle cerebral artery using trans-cranial Doppler²⁰⁴ and administration of sublingual NTG resulted in a mean $30 \pm 8\%$ increase in the human coronary artery²⁰⁵. The mean diameter of the MMA in healthy volunteers was 1,4 mm with an SD of 0,18 (pilot study). We therefore calculated that we would require 20 subjects to detect a difference of at least 10% in means at the 5% level of significance (power 90%).

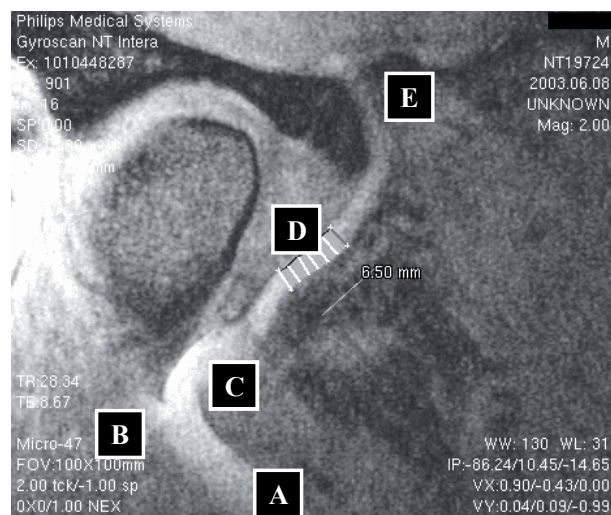


Figure 1 Anatomy of the MMA region and position of the measured segment. Explanation of letters: A= External Carotid Artery, B= Superficial Temporal Artery, C= Maxillary Artery, D= Middle Meningeal Artery, E= Foramen Spinosum

RESULTS

Three patients were excluded during the experiment. The first volunteer had an unexpected MRI finding, the second had a double extra cranial MMA (possibly an accessory meningeal artery) and in the third volunteer the MMA could not be reliably measured. In the remaining 19 subjects (9 males; mean age 21.8 ± 2.9 years) the MMA could be easily identified (Figure 1). The mean MMA diameter (extra cranial part) was

regel 1
regel 2
regel 3
regel 4
regel 5
regel 6
regel 7
regel 8
regel 9
regel 10
regel 11
regel 12
regel 13
regel 14
regel 15
regel 16
regel 17
regel 18
regel 19
regel 20
regel 21
regel 22
regel 23
regel 24
regel 25
regel 26
regel 27
regel 28
regel 29
regel 30
regel 31
regel 32
regel 33
regel 34
regel 35
regel 36
regel 37
regel 38
regel 39

1.5 ± 0.26 mm before and 1.79 mm ± 0.30mm after NTG administration (Table 1). The increase after NTG was 20.1% (CI: 12.9% to 27.3%; p<0.001) from baseline.

Table 1 MMA diameter at baseline and increase after sublingual nitroglycerin (NTG).

Subjects	N	Baseline (mm)	Post NTG (mm)	Difference post NTG vs baseline (mm)	
		Mean (SD)	Mean (SD)	Mean (SD)	% from baseline
All	19	1.50 (0.26)	1.79 (0.3)	0.29 (0.20)*	20.1%
Headache post NTG	11	1.52 (0.31)	1.87 (0.32)	0.34 (0.19)	23.9%
No headache post NTG	8	1.48 (0.18)	1.7 (0.27)	0.22 (0.20)	14.8%

(* = p<0.001)

Within five minutes after nitroglycerin administration, eleven volunteers experienced mild, bilateral, pulsating headache of short duration (<30 minutes) and without associated phonophobia, photophobia or nausea. None of the headaches fulfilled the IHS criteria for migraine. No adverse events or significant effects on blood pressure occurred. The mean diastolic blood pressure at baseline was 74.1 (SD 5.7) and the mean systolic blood pressure was 122.6 (SD 8.6). The mean MMA diameter in the 11 subjects who developed headache was 1.52 ± 0.31 mm before and 1.87 ± 0.32 mm after nitroglycerin administration as compared to 1.48 ± 0.18 mm before and 1.7 ± 0.27 mm after nitroglycerin administration in the 8 subjects who did not develop headache (CI for difference: -0.07 to 0.31; p=0.188; Table 1). The post-hoc power to detect a difference in MMA diameter increase of 9.1% between subjects with and without headache was only 27% (alpha 0.05, SD 0.18). Agreement between observers (intra class correlation) was 0.74 (0.7 or more is considered acceptable).

DISCUSSION

MRA in combination with a 47 mm microscopy coil is a novel, promising non-invasive method to study the MMA *in vivo*. The whole scan procedure takes 15 minutes making it very suitable for repeated clinical studies. Localization and measurement of the MMA was possible in 20 out of 22 subjects. The measurement precision of the used technique is 0.09 mm, which is sufficient for valid measurements of both the baseline MMA diameter as well as diameter changes after nitroglycerin administration²⁰⁶.

A relatively recent development in MRA is contrast-enhanced MRA (CE-MRA). For CE

MRA fast scan times and adequate timing based on a test bolus are required to avoid venous over projection of the jugular veins. After the injection of a test bolus, current available CE MRA methods acquire high contrast arterial signal in the first 10 seconds, within the time-window of arterial enhancement. Thereafter, the acquisition is continued to increase the resolution of the depicted arteries. With the injection of an intravenous contrast bolus of gadolinium the T1 of the blood is shortened and larger flip angles can be used to generate a stronger signal with improved background suppression and less signal saturation. CE MRA provides morphological information over a long track starting at the neck arteries via to the circle of Willis up to the distal intracranial smaller vessel segments. Extra-cranially, CE-MRA may also provide better resolution of the MMA. However, currently no studies have been performed in this matter. In spite of the advantages of CE-MRA, we used a non CE-MRA acquisition technique because of medical ethical concerns: in a CE-MRA protocol gadolinium contrast should be delivered twice in relatively short time (less than 30 minutes) with administration of nitroglycerin in between.

A potential limitation of this method may be that the observed diameter increase is overestimated due to increase of the blood flow velocity when using MRA (time of flight) diameter measurements. However, we do not think this is the case for two reasons. Firstly, Bednarczyk et al. measured an increase in global cerebral blood flow (positron emission tomography) after nitroglycerin administration without an increase in flow velocity in the middle cerebral artery (trans-cranial Doppler)²⁰⁷, and secondly, the contour of the blood vessel is automatically detected using MRI-MASS. An increase in flow velocity will increase the intravascular signal intensity, but this will probably not affect the automatic contour detection. A potential effect of flow velocity changes can however not be ruled out.

This new research method may have important implications for the study of migraine (notably for measuring the MMA during spontaneous and experimental migraine attacks and after treatment with antimigraine agents)²⁰⁸. The current study was not designed to prove or disprove a causal relationship between vasodilatation of the MMA and the occurrence of migraine headache. We found a mean 23.9% dilatation of the MMA in subjects with non-migrainous headache after nitroglycerin administration and 14.8% dilatation in those without headache. This difference was non-significant which may have been due to the small number of study subjects. The post hoc power to detect a statistically significant difference in vasodilatation between subjects with and without headache was only 27%. Further studies are needed to address this issue. Besides migraine, this method might be of interest for other neurovascular research areas, such as meningeal vasospasms in subarachnoid hemorrhage²⁰⁹.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

— |

| —

—————

— |

| —

CHAPTER 6

CEREBRAL BLOOD FLOW RESPONSE TO NITROGLYCERIN PREDICTS THE OCCURRENCE OF A PROVOKED MIGRAINE ATTACK

Submitted



ABSTRACT

Background

Nitroglycerin (NTG) triggers migraine attacks in susceptible migraine patients. The mechanism of action is unclear. The cerebrovascular response to NTG may be impaired in migraine patients, however, previous studies are inconclusive. In this study we assessed the cerebrovascular response to NTG in healthy volunteers and in migraine patients with and without a provoked attack.

Methods

In a double blind desing migraine patients (n=32) received NTG (n=27) IV 0.5 µg/kg/min for 20 min or placebo (n=5). Healthy volunteers (n=12) all received NTG. Using 3T MRA, we measured blood flow and diameter in the internal carotid arteries (ICA) and basilar artery (BA) as well as diameters of the middle meningeal (MMA), external carotid (ECA), middle cerebral (MCA) and posterior cerebral (PCA) arteries at baseline and during infusion of NTG.

Results

During infusion of nitroglycerin, ICA blood flow decreased 118.9 ml/min in healthy volunteers and 19.5 ml/min in migraine patients (p=0.05). A sub-analysis of migraine patients showed an ICA blood flow decrease of 100.4 ml/min in patients who did not develop a provoked migraine attack after NTG (n=7) compared to an increase of 10.2 ml/min in patients who developed a provoked migraine attack (n=20; p=0.01). Blood flow in the BA did not change. Diameters of all selected blood vessels increased significantly (p<0.01) during infusion of NTG without differences between groups.

Conclusions

The cerebral blood flow response to nitroglycerin is impaired in migraine patients mainly due to an impaired response in patient who develop a migraine attack after approximately 4 hours.

INTRODUCTION

Migraine is a severe paroxysmal neurovascular disorder⁷⁵. Attacks can be reliably^{55,78} and reproducibly⁵⁶ triggered in migraine susceptible subjects using the vasoactive drug nitroglycerin. In healthy volunteers without a family history of migraine it is very seldom that migraine attacks are triggered after infusion of NTG¹¹⁶. The mechanism of nitroglycerin in migraine is unclear²¹⁰. Nitroglycerin is an exogenous donor of nitric oxide¹⁰⁶, which is involved in central pain mechanism¹⁰⁷ and regulation of cerebral blood flow¹⁰⁸. Infusion of NTG has shown to increase the diameter of the middle cerebral artery¹⁰⁹ and meningeal media artery¹¹⁰ as well as to decrease blood flow velocity in the internal carotid artery and middle cerebral artery¹¹¹⁻¹¹³. The effects of NTG on cerebral blood flow are caused either through the release of calcitonin gene related peptide (CGRP) from the trigeminal nerve^{114,115} or via a direct effect on vascular smooth muscle cells in blood vessels¹⁰⁶. Whether there is a difference in cerebrovascular response to NTG between migraine patients and healthy controls is unclear. One study suggested an increased cerebrovascular response during NTG infusion in migraine patients¹¹⁷, whereas in a second study no increased response was observed.¹¹⁸

Infusion of NTG triggers a migraine attack in approximately 60 to 80% of migraine patients^{55,56,78,116,124,125}. Why some patients are susceptible to NTG and others not is unknown. Migraine patients without aura are more susceptible to NTG as compared to patients with aura⁵⁵. Furthermore, in a NTG provocation study an association between increase in plasma CGRP and the provocation of a migraine attack has been found¹²⁴. Whether the cerebrovascular response to NTG is different in patients who develop a provoked attack as compared to patients without an attack is unclear.

The primary aim of this study was to assess the cerebrovascular response (blood flow and blood vessel diameters) to infusion of NTG in healthy volunteers and migraine patients. The hypothesis is that the cerebrovascular response to nitroglycerin is impaired in migraine patients compared to healthy volunteers. The secondary aim is to assess the relation between cerebrovascular response to NTG and the development of a provoked migraine attack.

____ regel 1
____ regel 2
____ regel 3
____ regel 4
____ regel 5
____ regel 6
____ regel 7
____ regel 8
____ regel 9
____ regel 10
____ regel 11
____ regel 12
____ regel 13
____ regel 14
____ regel 15
____ regel 16
____ regel 17
____ regel 18
____ regel 19
____ regel 20
____ regel 21
____ regel 22
____ regel 23
____ regel 24
____ regel 25
____ regel 26
____ regel 27
____ regel 28
____ regel 29
____ regel 30
____ regel 31
____ regel 32
____ regel 33
____ regel 34
____ regel 35
____ regel 36
____ regel 37
____ regel 38
____ regel 39

METHODS

Subjects

In total 32 migraine patients (without aura n=27 and with aura n=5) and 12 healthy volunteers were included. Patients were recruited from the neurology outpatient clinic of Leiden University Medical Centre. Inclusion criteria for migraine patients were (1) age between 18 and 55 years, (2) diagnosis of migraine according to the criteria of the IHS³, (3) baseline attack frequency between 1 attack per 2 months to 4 attacks per month in the six months prior to the study, (4) moderate or severe headache during spontaneous migraine attacks. Exclusion criteria were (1) more than 10 days of headache per month, (2) inability to differentiate between migraine and other forms of headache, (3) contra-indications for use of triptans, (4) current use of vasoactive drugs and (5) MRI specific contra-indications (such as claustrophobia). Healthy volunteers were recruited among hospital staff, medical students and relatives of migraine patients. Inclusion criteria for healthy volunteers were (1) age between 18 and 55 years. Exclusion criteria were (1) personal or first degree relative history of migraine, (2) headache on more than 2 days per month, (3) MRI specific contra-indications and (4) current use of vasoactive drugs. The study was approved by the local medical ethical committee and the subjects gave informed consent prior to the start of the study.

Experimental procedure

All subjects arrived at the hospital without headache between 8 and 10 a.m. on the day of the study. No medication, coffee, tea or alcohol was allowed 12 hours prior to the start of the experiment. In migraine patients the last spontaneous migraine attack was at minimum three days prior to the experiment. Healthy volunteers were scanned at baseline and during infusion of NTG 0.5 µg/kg/min during 20 min (open label). Migraine patients were scanned at baseline, during infusion of NTG/ placebo (double blind) and during a provoked migraine attack (or 6 hours after infusion of NTG/placebo). Duration of scan sessions was approximately 25 minutes. Between the baseline session and the NTG/placebo session patients remained in the scanner and the NTG/placebo session started 10 minutes after the start of the infusion. Heart rate and blood pressure were monitored before and after the MR session. Two days after the experiment subjects were contacted by telephone to make sure no migraine attack had occurred within 12 hours after the experiment.

Magnetic resonance angiography

The MR investigations were performed on a 3.0-Tesla whole-body system (Philips

Medical Systems, The Netherlands). The MRA protocol consisted of two parts. Part A) Diameter protocol: a thick two-dimensional phase contrast (2D PC) sagittal localiser survey through the circle of Willis, followed by a three-dimensional time-of-flight (3D TOF) MRA sequence to visualise the BA and ECA, ICA, and MCA on both sides This scan had the following imaging parameters: repetition time / echo time (TR/TE): 22 ms / 3.5 ms; flip angle 15°; field of view: 220 x 220 mm; number of excitations: 1; slice orientation: transverse; slice thickness: 0.65 mm; number of slices: 200; scan percentage 100%, matrix reconstruction size: 512 x 512 resulting in a nominal voxel size (x,y,z) of 0.43 x 0.43 x 0.65 mm; total acquisition time: 4min 30sec. Based on the reconstruction of this 3D-TOF a second 3D-TOF with a higher spatial resolution was performed to visualise the extra- and intracranial parts of the MMA on both sides. This scan had the following imaging parameters: TR/TE: 15 ms / 2.1 ms; flip angle 15°; field of view: 200 x 200 mm; number of excitations: 1; slice orientation: transverse; slice thickness: 0.25 mm; number of slices: 130; scan percentage 100%, matrix reconstruction size: 512 x 512 resulting in a nominal voxel size (x,y,z) of 0.39 x 0.39 x 0.25 mm; total acquisition time: 8min 31sec.

Part B) Blood flow protocol (in BA and ICA): On the basis of two thick slab localizer MRA scans in the coronal and sagittal plane, a 2-dimensional phase-contrast (2D-PC) section was positioned perpendicular on the ICAs and the BA at the level of the skull base to measure the volume flow. The MRA volume flow measurements in the present study are derived from previously developed and optimized protocols²¹¹⁻²¹⁴. Acquisition parameters: repetition time / echo time (TR/TE): 16 ms / 8.5 ms; flip angle 10°; field of view: 150 x 150 mm; number of excitations: 20; slice orientation: transverse; slice thickness: 5.0 mm; number of slices: 1; scan percentage 100%; PC velocity encoding: 140 cm/s; matrix reconstruction size: 256 x 256 resulting in a nominal voxel size (x,y,z) of 0.59 x 0.59 x 50 mm; total acquisition time: 56sec. Figure 1 illustrates the positioning of the 2D PC section through the internal carotid arteries (ICAs) and the basilar artery (BA). On an independent workstation, quantitative flow values were calculated in each vessel by integrating across manually drawn regions of interest that enclosed the vessel lumen closely.

Image post processing: diameter calculations

All MRA images were transferred to a independent workstation for quantitative analysis using the QMRA software package developed at our institution. A full description of the contour detection methods used and the validation have been described previously²¹⁵. The software provides automated contour detection and quantification of the luminal

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

boundaries in selected vessel segments in 3D MRA datasets. The only user interaction required is the definition of the vessel segment of interest by placing a proximal and distal point in the 3D dataset. Subsequently, the software detects a 3D path line following the centre of the vessel lumen and cross-sectional MPR's are generated perpendicular to the centreline at 0.5 mm intervals. In each of these MPR's a contour around the vessel lumen is detected automatically. From these contours, based on the assumption of circular vessel cross-sections, the average diameter of the selected vessel segment is derived. Blood vessel segments were selected as following: A) The MMA was measured in an extra cranial segment (start at the origo in the maxillary artery and stop 5 to 6 mm distal). B) The ECA start at the origo of the superficial temporal artery and stop 10 mm proximal. C) The ICA, start just proximal of the Syphon and stop 15 mm proximal. D) The MCA, start after A1 segment and stop 8 mm distal. E) The BA, start origo posterior cerebral artery stop 12 mm proximal. F) The PCA, start at the origo in the basilar artery and stop 8 mm distal). Location of measured vessel segments were kept constant within subjects.

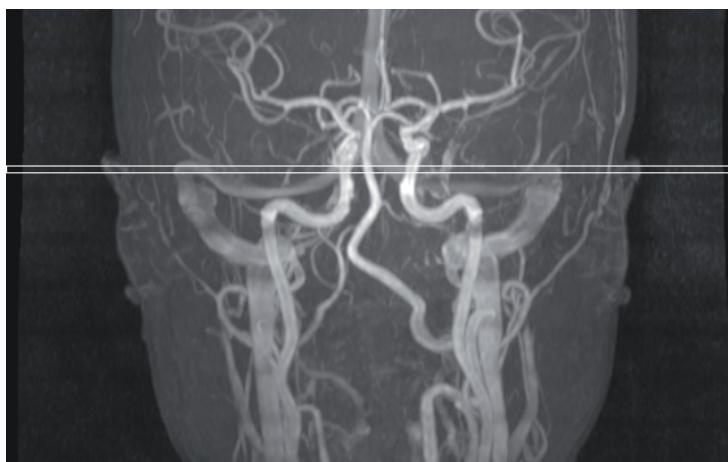


Figure 1 Magnetic resonance angiography, coronal maximum intensity projection. Horizontal line indicates the positioning of the 2-dimensional phase-contrast section through the ICA and the BA.

Statistical analysis

Data were analysed using SPSS 12.0.1 (SPSS Inc, Chicago, USA). Left-to-right differences in diameters for bilateral blood vessels (MMA, ICA, ECA, MCA and PCA) were tested using the paired t-test. Since differences were not significant (as shown in the results section) means of the right and left vessel were used throughout this paper. Difference in blood vessel diameter and blood flow at baseline between migraine patients and

healthy volunteers was tested using the Mann Whitney U test. Linear mixed models were used to analyse the effect of nitroglycerin, subjects (healthy volunteers/ migraine) and provoked migraine attack (yes/ no) on vessel diameters and blood flow. Data from patients receiving placebo was not used for statistical testing. $P < 0.05$ was considered significant.

RESULTS

In total 32 migraine patients (27 NTG and 5 Placebo) and 12 healthy volunteers were included in the study (Table 1). A provoked migraine attack was observed in 20 migraine patients (74%) and no attack after placebo or in healthy volunteers. The median time to attack was 3.75 hours (range 1.5 – 5.5 hours).

Table 1 Demographic characteristics of study participants

Intervention	HV (n=12)		Migraine (n=32)	
	NTG		NTG (27)	Placebo (n=5)
Attack	No	Yes (n=20)	No (n=7)	No
Age in years (SD)	40.6 (11.1)	45.5 (8.5)	34 (8.9)	44.8 (13.3)
Ratio female to males	10 : 2	15 : 5	7 : 0	3 : 2
Ratio MO to MA		17 : 3	6 : 1	4 : 1
Attack frequency; mean (SD)		2.6 (1.0)	2.1 (0.38)	2.4 (1.1)

HV denotes healthy volunteer, MO migraine without aura, MA migraine with aura.

Baseline measurements

There were no differences in blood vessel diameters or blood flow at baseline between migraine patients and healthy volunteers (Table 2 and 4).

Side to side differences of blood vessel diameter

There were no ($p > 0.05$) right to left differences for the diameters of the five bilateral blood vessels (MMA, ICA, ECA, MCA, PCA) in any of the conditions. Therefore, mean right-left diameters are presented throughout.

Blood flow changes in BA and ICA during NTG

Blood flow in the ICA decreased 118.9 ml/min in healthy controls and decreased 19.5 ml/min in migraine patients ($p = 0.05$; Table 2). In migraine patients who later developed a migraine attack blood flow in the ICA increased 7.2 ml/min during NTG and decreased

112.5 ml/min in patients without an attack ($p=0.01$; Table 3). Blood flow changes in the BA during NTG was not different between migraine patients and healthy volunteers nor in migraine patients with and without an attack.

Table 2 Blood flow in BA, ICA and total cerebral blood flow in migraine patients and healthy volunteers at baseline and during infusion of nitroglycerin or placebo.

Blood vessel	Subjects	Inter- vention	N	A)	B)	Change (B vs A) ml/min (%)
				Baseline ml/min (SD)	During NTG or placebo ml/min (SD)	
BA	HV	NTG	12	145.8 (55.5)	149.0 (50.9)	3.3 (2.3)
		NTG	27	174.6 (68.7)	169.3 (57.9)	-5.4 (-3.1)
	Placebo	5	170.5 (39.4)	128.6 (49.9)	-41.9 (-24.6)	
ICA	HV	NTG	12	700.5 (200.4)	581.7 (154.8)	-118.9 (-17.0)*
		NTG	27	577.1 (121.6)	557.5 (139.4)	-19.5 (-3.4)
	Placebo	5	542.0 (211.1)	523.2 (161.9)	-18.8 (-3.5)	
TCBF	HV	NTG	12	850.9 (199.8)	735.1 (139.9)	-115.9 (-13.6)
		NTG	27	751.7 (117.3)	726.8 (149.5)	-24.9 (-3.3)
	Placebo	5	712.6 (202.4)	651.8 (198.8)	-60.7 (-8.5)	

HV denotes healthy volunteers, NTG nitroglycerin, ICA internal carotid artery, BA basilar artery, tCBF total cerebral blood flow. *ICA blood flow is significantly different between healthy volunteers and migraine ($p=0.05$)

Table 3 Blood flow in BA, ICA and total cerebral blood flow in migraine patients with and without a provoked migraine attack at baseline and during infusion of nitroglycerin.

Blood vessel	Subjects	Inter- vention	Migraine attack	N	A)	B)	Change (B vs A) ml/min (%)
					Baseline ml/min (SD)	During NTG or placebo ml/min (SD)	
BA	Migraine	NTG	Yes	20	173.7 (69.4)	170.8 (64.7)	-2.8 (-1.6)
			No	7	177.2 (71.9)	165.0 (36.8)	-12.2 (-6.9)
ICA	Migraine	NTG	Yes	20	589.6 (128.5)	599.9 (128.1)	10.2 (1.7)*
			No	7	542.9 (101.2)	442.5 (103.3)	-100.4 (-18.5)
TCBF	Migraine	NTG	Yes	20	763.3 (124.1)	770.7 (140.7)	7.4 (1.0)
			No	7	720.1 (97.7)	607.5 (105.4)	-112.5 (-15.6)

HV denotes healthy volunteers, NTG nitroglycerin, ICA internal carotid artery, BA basilar artery, tCBF total cerebral blood flow. *ICA blood flow is significantly different between healthy volunteers and migraine ($p=0.05$)

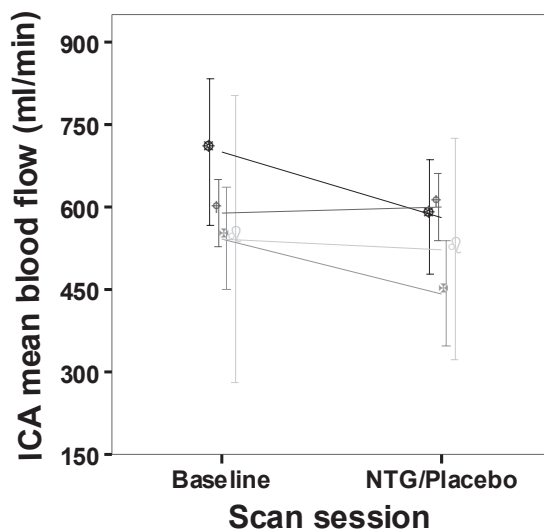


Figure 2A Blood flow in internal carotid artery at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

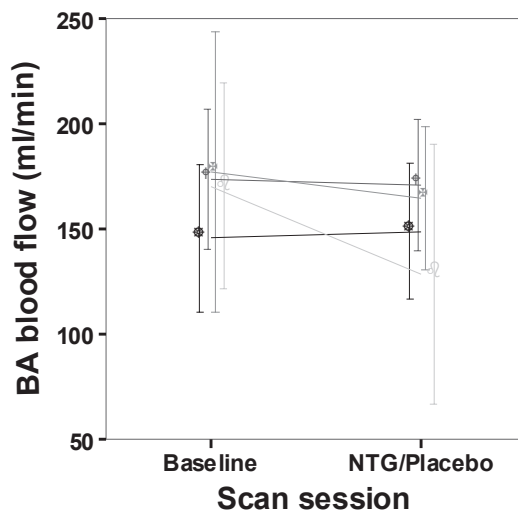


Figure 2B Blood flow in basilar artery at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

- regel 1
- regel 2
- regel 3
- regel 4
- regel 5
- regel 6
- regel 7
- regel 8
- regel 9
- regel 10
- regel 11
- regel 12
- regel 13
- regel 14
- regel 15
- regel 16
- regel 17
- regel 18
- regel 19
- regel 20
- regel 21
- regel 22
- regel 23
- regel 24
- regel 25
- regel 26
- regel 27
- regel 28
- regel 29
- regel 30
- regel 31
- regel 32
- regel 33
- regel 34
- regel 35
- regel 36
- regel 37
- regel 38
- regel 39

Table 4 Blood vessel diameters of six selected cranial blood vessel in healthy volunteers and migraine patients at baseline and during infusion of nitroglycerin or placebo.

Blood vessel	Subjects	Inter-vention	N	A)	B)	Change (B vs A)
				Baseline	During NTG or placebo	
				mm (SD)	mm (SD)	mm (% from A)
MMA	HV	NTG	12	1.61 (0.11)	1.93 (0.21)	0.32 (19.9)*
	Migraine	NTG	27	1.65 (0.18)	1.93 (0.24)	0.27 (16.4)*
	Migraine	Placebo	5	1.67 (0.73)	1.64 (0.12)	-0.02 (-1.2)
ECA	HV	NTG	12	3.63 (0.46)	4.61(0.39)	0.98 (27.0)*
	Migraine	NTG	27	3.46 (0.38)	4.50 (0.38)	1.05 (30.3)*
	Migraine	Placebo	5	3.51 (0.27)	3.56 (0.39)	0.05 (1.4)
ICA	HV	NTG	12	4,87 (0.16)	5,38 (0.30)	0.51 (10.5)*
	Migraine	NTG	27	4.81 (0.49)	5.32 (0.42)	0.51 (10.6)*
	Migraine	Placebo	5	4,86 (0.41)	5,02 (0.44)	0.15 (3.1)
MCA	HV	NTG	12	3,14 (0.15)	3,46 (0.24)	0.33 (10.5)*
	Migraine	NTG	27	3.16 (0.29)	3.52 (0.24)	0.37 (11.7)*
	Migraine	Placebo	5	3,10 (0.20)	3,10 (0.22)	-0.01 (-0.3)
BA	HV	NTG	12	3,00 (0.42)	3,41 (0.50)	0.41 (13.7)*
	Migraine	NTG	27	2.95 (0.53)	3.57 (0.57)	0.61 (20.7)*
	Migraine	Placebo	5	2,86 (0.42)	2,80 (0.38)	-0.06 (-2.1)
PCA	HV	NTG	12	2.53 (0.16)	2.68 (0.11)	0.14 (5.5)*
	Migraine	NTG	27	2.55 (0.15)	2.72 (0.19)	0.17 (6.7)*
	Migraine	Placebo	5	2.67 (0.15)	2.62 (0.28)	-0.04 (-1.5)

HV denotes healthy volunteers, NTG nitroglycerin, MMA middle meningeal artery, ECA external carotid artery, ICA internal carotid artery, MCA middle cerebral artery, BA basilar artery and PCA posterior cerebral artery. *Significant increase in diameters in all blood vessels during infusion of NTG as compared to baseline ($p < 0.01$).

Diameter changes during infusion of NTG

Compared to baseline NTG caused a significant vasodilatation of all six selected blood vessels (Table 4 and Figures 3A to F; $p < 0.01$ for all blood vessels). The immediate NTG-induced diameter increase was larger in the extra-cerebral blood vessels (MMA and ECA), with an increase ranging from 16.4% to 30.3% as compared to the diameter increase seen in the intra-cranial blood vessels (ICA, MCA, BA and PCA), with an

increase ranging from 5.5% to 20.7%. The diameter increase during NTG was not significantly different ($p > 0.05$) between migraine patients and healthy volunteers (Table 4) nor between migraine patients with a provoked attack compared to patients without an attack (Table 5) in all measured blood vessels.

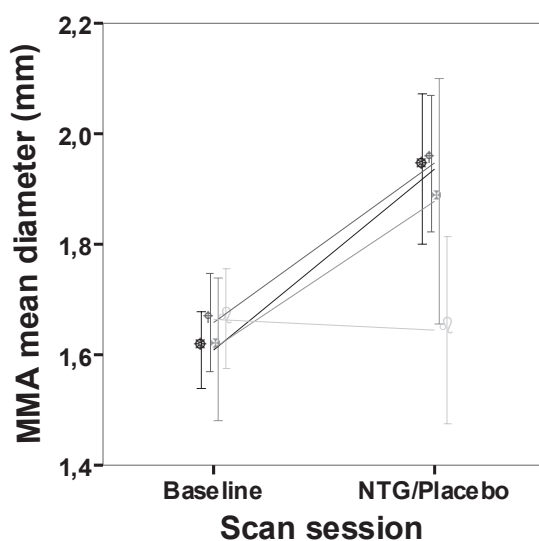


Figure 3A Blood vessel diameter of the MMA at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

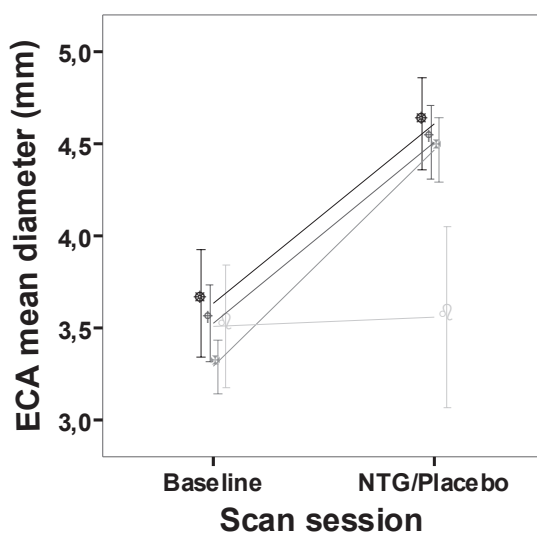


Figure 3B Blood vessel diameter of the ECA at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

regel 1
 regel 2
 regel 3
 regel 4
 regel 5
 regel 6
 regel 7
 regel 8
 regel 9
 regel 10
 regel 11
 regel 12
 regel 13
 regel 14
 regel 15
 regel 16
 regel 17
 regel 18
 regel 19
 regel 20
 regel 21
 regel 22
 regel 23
 regel 24
 regel 25
 regel 26
 regel 27
 regel 28
 regel 29
 regel 30
 regel 31
 regel 32
 regel 33
 regel 34
 regel 35
 regel 36
 regel 37
 regel 38
 regel 39

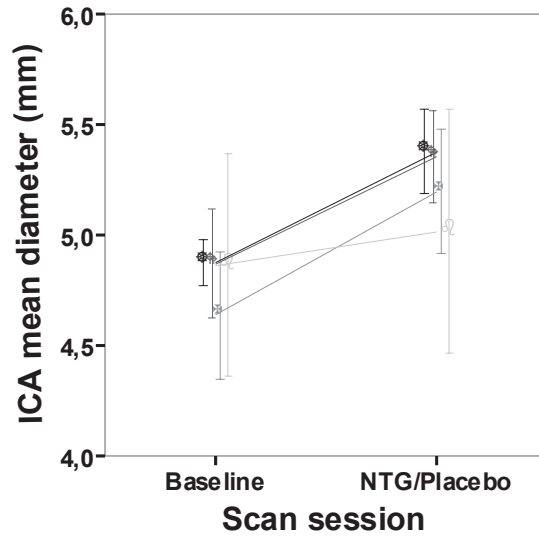


Figure 3C Blood vessel diameter of ICA at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

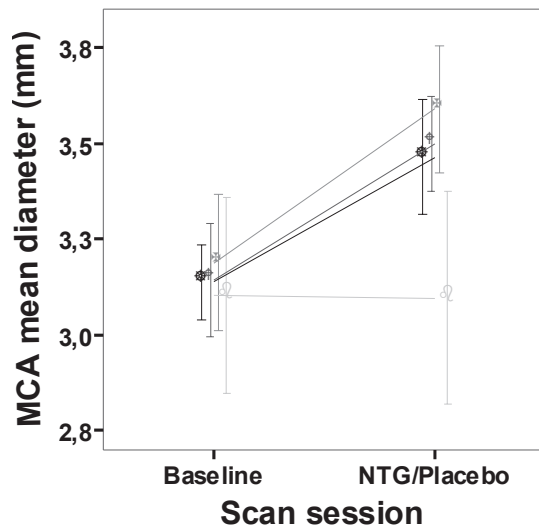


Figure 3D Blood vessel diameter of MCA at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

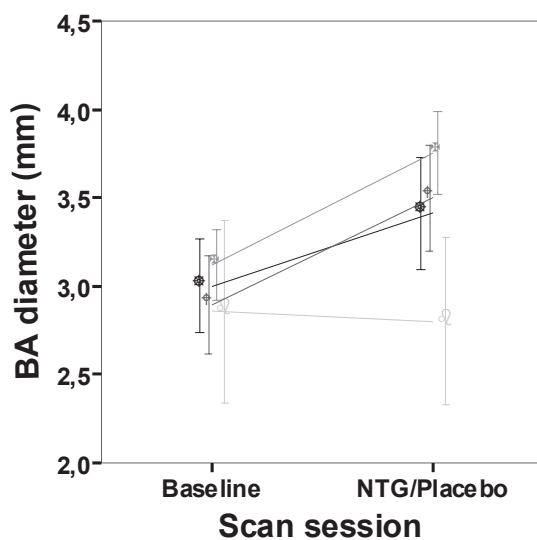


Figure 3E Blood vessel diameter of BA at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

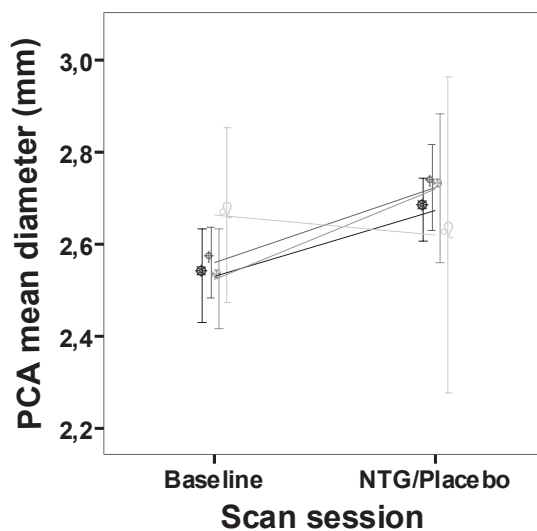


Figure 3F Blood vessel diameters in selected blood vessels at baseline and during infusion of nitroglycerin or placebo. (■ Healthy volunteers (NTG), ● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

- regel 1
- regel 2
- regel 3
- regel 4
- regel 5
- regel 6
- regel 7
- regel 8
- regel 9
- regel 10
- regel 11
- regel 12
- regel 13
- regel 14
- regel 15
- regel 16
- regel 17
- regel 18
- regel 19
- regel 20
- regel 21
- regel 22
- regel 23
- regel 24
- regel 25
- regel 26
- regel 27
- regel 28
- regel 29
- regel 30
- regel 31
- regel 32
- regel 33
- regel 34
- regel 35
- regel 36
- regel 37
- regel 38
- regel 39

Table 5 Blood vessel diameters of six selected cranial blood vessel in migraine patients with and without a provoked migraine attack at baseline and during infusion of nitroglycerin.

Blood vessel	Subjects	Intervention	Migraine attack	N	A)	B)	Change (B vs A) mm (% from A)
					Baseline mm (SD)	During NTG or placebo mm (SD)	
MMA	Migraine	NTG	Yes	20	1.66 (0.19)	1.95 (0.24)	0.28 (16.9)
	Migraine	NTG	No	7	1.61 (0.12)	1.88 (0.20)	0.27 (16.8)
ECA	Migraine	NTG	Yes	20	3.53 (0.42)	4.51 (0.43)	1.00 (28.3)
	Migraine	NTG	No	7	3.29 (0.16)	4.47 (0.19)	1.18 (35.9)
ICA	Migraine	NTG	Yes	20	4,87 (0.53)	5,36 (0.45)	0.49 (10.1)
	Migraine	NTG	No	7	4,64 (0.31)	5,20 (0.31)	0.56 (12.1)
MCA	Migraine	NTG	Yes	20	3,14 (0.32)	3,50 (0.26)	0.36 (11.5)
	Migraine	NTG	No	7	3,19 (0.19)	3,59 (0.18)	0.4 (12.5)
BA	Migraine	NTG	Yes	20	2,89 (0.60)	3,50 (0.64)	0.6 (20.8)
	Migraine	NTG	No	7	3,12 (0.21)	3,75 (0.25)	0.63 (20.2)
PCA	Migraine	NTG	Yes	20	2.56 (0.16)	2.73 (0.20)	0.17 (6.6)
	Migraine	NTG	No	7	2.53 (0.12)	2.72 (0.17)	0.20 (7.9)

HV denotes healthy volunteers, NTG nitroglycerin, MMA middle meningeal artery, ECA external carotid artery, ICA internal carotid artery, MCA middle cerebral artery, BA basilar artery and PCA posterior cerebral artery.

DISCUSSION

In the present study we found that the decrease in ICA blood flow during NTG infusion was more pronounced in healthy volunteers as compared to migraine patients. Changes in BA blood flow and blood vessel diameter were not different between groups. Previous NTG studies in migraine showed either no difference in blood velocity decrease in the middle cerebral artery between migraine and controls¹¹⁸, or a more pronounced decrease in migraine patients¹¹⁷. An explanation for the difference in results could be that these studies did not take into account the occurrence of a provoked attack. In this study the difference between migraine and healthy volunteers is mainly explained by a difference between migraine patients with and without a provoked attack. Patient without an attack showed a decrease in ICA blood flow similar to healthy volunteers whereas in patients with an attack ICA blood flow did not decrease.

Blood flow in the ICA is affected by several parameters; i) ICA blood vessel diameter, ii) cardiac output and iii) vasomotor tone in small resistance vessels. The ICA diameter increased during NTG infusion, but there was no difference between groups. Nitroglycerin has shown to decrease cardiac output²¹⁶. In this study we did not measure cardiac output but we did not observe a difference in blood pressure response during NTG infusion between groups (data not shown), suggesting that there was no difference in decrease of cardiac output between groups. So a difference in vasomotor tone of small resistance vessels might be the main explanation for the observed difference between patients with and without an attack.

Many factors are involved in the regulation of cerebral blood flow; for review see Hamel²¹⁷. Nitroglycerin has shown to affect cerebral blood flow via release of CGRP from trigeminal perivascular nerves^{114,115} and through a direct effect on vascular smooth muscle cells¹⁰⁶. An increased release of CGRP during NTG infusion in patients with an attack would fit previous findings in provoked attacks: the occurrence of a provoked attack was associated with an increase in CGRP during NTG provocation¹²⁴. Whether there could be a different effect of NTG on vascular smooth muscle cells between migraine patients with an attack as compared to patients without an attack is unclear.

Another interesting finding was that nitroglycerin decreased cerebral blood flow in healthy volunteers, whereas other vasodilators have shown to increase tCBF in studies using phase contrast MRA. Acetazolamide infusion showed a tCBF increase of 41% in healthy volunteers²¹⁸ and CO₂ increased tCBF by 64%²¹⁹. An explanation could be that cardiac output is increased in acetazolamide²²⁰ and decreased during nitroglycerin infusion²¹⁶. Future studies on the effect of nitroglycerin on cerebral blood flow should include measures for cardiac output.

In conclusion, the ICA blood flow response to nitroglycerin is impaired in migraine patients compared to healthy volunteers, mainly due to an impaired response in patients who developed a provoked migraine attack after several hours. These findings suggest that provocation of an attack after NTG is associated with an impaired response in small resistance blood vessels.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

— |

| —

—————

— |

| —

CHAPTER 7

MIGRAINE HEADACHE IS NOT ASSOCIATED WITH CEREBRAL OR MENINGEAL VASODILATATION - A 3T MAGNETIC RESONANCE ANGIOGRAPHY STUDY

Brain. 2008 May 23 (epub ahead of print)



ABSTRACT

Background

Migraine headache is widely believed to be associated with cerebral or meningeal vasodilatation. Human evidence for this hypothesis is lacking. 3 Tesla Magnetic resonance angiography (3T MRA) allows for repetitive, non-invasive, sensitive assessment of intracranial vasodilatation and blood flow. Nitroglycerine (NTG) can faithfully induce migraine attacks facilitating pathophysiological studies in migraine.

Methods

Migraineurs (n=32) randomly received NTG (IV 0.5 µg/kg/min for 20 min; n=27) or placebo (n=5; for blinding reasons). Using 3T MRA, we measured: a) blood flow in the basilar (BA) and internal carotid (ICA) arteries and b) diameters of the middle meningeal (MMA), external carotid (ECA), ICA, middle cerebral (MCA), BA and posterior cerebral (PCA) arteries at three timepoints: i) at baseline, outside an attack; ii) during infusion of NTG or placebo; and iii) during a provoked attack or, if no attack had occurred, at 6 hours after infusion.

Findings

Migraine headache was provoked in 20/27 (74%) migraineurs who received NTG, but in none of the five patients who received placebo. The headache occurred between 1.5 – 5.5 hrs after infusion and was unilateral in 18/20 (90%) responders. During NTG (but not placebo) infusion, there was a transient 6.7% – 30.3% vasodilatation ($p < 0.01$) of all blood vessels. During migraine, blood vessel diameters were no different from baseline, nor between headache and non-headache sides. There were no changes in BA and ICA blood flow during either NTG infusion or migraine.

Interpretation

In contrast to widespread belief, migraine attacks are not associated with vasodilatation of cerebral or meningeal blood vessels. Future antimigraine drugs may not require vasoconstrictor action.

INTRODUCTION

Migraine is a neurovascular disorder typically characterised by attacks of severe, throbbing, unilateral headache, associated autonomic symptoms, and, in one third of patients, focal neurological aura symptoms ⁷⁵. Since the seminal work by Wolff and colleagues ¹⁹⁸, showing that stimulation of cerebral and meningeal arteries caused headache, there is a widespread belief that vasodilatation of intracranial blood vessels is the underlying mechanism for migraine headache ²²¹. This hypothesis was further fed by a number of other observations. Balloon dilatation of the middle cerebral artery (MCA) may cause migraine-like headache ²²². Vasoactive substances such as the nitric oxide (NO) donor nitroglycerin (NTG) ⁷⁸ and calcitonin gene related peptide (CGRP) ⁶² can trigger migraine in susceptible subjects. In fact, the recent development of novel CGRP antagonists for treating migraine attacks was at least partly based on the hypothesis that prevention or reversal of vasodilation would block migraine headache ^{223,224}. Animal and *in situ* pharmacological experiments ^{75,225} and human *in vivo* studies using transcranial Doppler (TCD) ^{204,226,227} have shown that acute antimigraine agents (ergots and triptans) constrict cerebral and meningeal blood vessels ²²⁸. In fact, the triptan class was specifically designed to selectively constrict intracranial blood vessels ²²¹.

The role of vasodilatation in migraine has been vividly debated in the past (for review see: ²²⁹) and more recently ^{75,87}. Some researchers view vasodilation of meningeal or cerebral blood vessels as a primary trigger for migraine headaches, and consider vasoconstriction necessary for acute antimigraine efficacy ²³⁰. Others feel that vasodilation is a secondary phenomenon, due to activation of the trigeminovascular system and release of vasoactive neuropeptides. Vasodilation would primarily be involved in sustaining and worsening of the headache during migraine attacks ⁷⁹. A third line of thinking holds that vasodilation is irrelevant or, at best, "an innocent bystander" in the pathogenesis of migraine headache. Consequently, vasoconstriction may not be needed to treat migraine headaches ^{231 232,233}. This would be an enormous advantage as the currently available most effective antimigraine agents, triptans and ergots, all possess (sometimes strong and sustained) vasoconstrictor activity ²³⁴. They may cause myocardial and cerebral ischaemia in patients with (risk factors for) vascular disease ²³⁵. Novel antimigraine agents, which are devoid of vasoconstrictor activity, would be safer and could thus also be used by the many migraineurs with vascular disease.

Remarkably, the three opposing views on the role of vasodilation in migraine are all primarily based on extrapolations of observations in experimental animal models, with

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____ very little evidence from human studies. This is primarily due to lack, until recently,
regel 2 _____ of sensitive non-invasive imaging techniques to directly and reliably assess intracranial
regel 3 _____ blood flow and blood vessel diameters in humans. Previous studies have used invasive
regel 4 _____ methods such carotid angiography ²³⁶, or could only indirectly estimate diameter changes
regel 5 _____ of cerebral blood vessels using TCD ^{204,237}. Meningeal blood vessels proved too small to be
regel 6 _____ investigated quantitatively. With the advent of 3 Tesla Magnetic Resonance Imaging (3T
regel 7 _____ MRA) a sensitive and non-invasive imaging technique has become available to reliably
regel 8 _____ measure intracranial blood flow and diameter changes of cerebral and meningeal blood
regel 9 _____ vessels ²³⁸ as small as the middle meningeal artery (MMA) ¹¹⁰.

regel 10 _____
regel 11 _____ Infusion of NTG can reliably and faithfully provoke migraine headaches in migraineurs
regel 12 _____ ^{55,83,210}. The response to NTG infusion is typically biphasic: an initial, brief and mild bilateral
regel 13 _____ headache during the infusion in nearly all migraine and non-migraine study subjects ⁸³,
regel 14 _____ followed by a typical migraine, 4 to 5 hours later, in 60% to 80% of migraine. but
regel 15 _____ not in non-migraine study subjects. ^{55,78} The symptomatology of provoked attacks is
regel 16 _____ no different from that of spontaneous attacks of migraine without aura ⁷⁸, including
regel 17 _____ premonitory symptoms ⁵⁶, response to anti-migraine drugs ²³⁹, and increase of CGRP, a
regel 18 _____ marker for activation of the trigeminovascular system ¹²⁴. This provocation model has
regel 19 _____ greatly facilitated the logistics of studying pathophysiological changes during migraine
regel 20 _____ attacks.

regel 21 _____
regel 22 _____ In the present study we used 3T MRA to intra-individually compare: a) blood flow in
regel 23 _____ the basilar (BA) and internal carotid (ICA) arteries; and b) the diameters of the external
regel 24 _____ carotid (ECA), internal carotid (ICA), middle cerebral (MCA), BA, posterior cerebral
regel 25 _____ (PCA) and middle meningeal arteries (MMA) between three conditions: i) at baseline,
regel 26 _____ outside an attack; ii) during infusion of NTG or placebo (to assess the immediate vascular
regel 27 _____ effects of NTG); and iii) during NTG-provoked migraine attacks or, if no attack had
regel 28 _____ occurred, at 6 hours post infusion (to assess whether migraine attacks are associated
regel 29 _____ with vasodilatation). We will demonstrate that there is no detectable vasodilatation of
regel 30 _____ cerebral or meningeal blood vessels during NTG-provoked migraine attacks, suggesting
regel 31 _____ that vasoconstriction may not be required to treat migraine headaches.

METHODS

Subjects

In total 32 migraine patients (n = 5 with aura; n = 27 without aura) were recruited from the neurology outpatient clinic of Leiden University Medical Centre. Inclusion criteria were: i) age between 18 and 55 years; ii) diagnosis of migraine according to the diagnostic criteria of the International Headache Society ³; iii) an average attack frequency between 1 - 8 attacks per 2 months in the six months prior to the study; and iv) moderate or severe headache during spontaneous migraine attacks. Exclusion criteria included: i) more than 10 days of headache per month; ii) inability to differentiate between migraine and other forms of headache; iii) contra-indications for the use of triptans; iv) current use of vasoactive drugs; and v) MRI-specific contra-indications (such as claustrophobia). The study was approved by the local medical ethics committee and the subjects gave informed consent prior to the start of the study.

Experimental procedure and NTG provocation

All subjects arrived at the hospital between 8 and 10 a.m. on the day of the study. No medication, coffee, tea or alcohol was allowed in the 12 hours prior to the start of the experiment. From one hour before the experiments until the very end of the experiments, study subjects were not allowed to smoke. Patients had to be free of migraine for at least the three days prior to the study day and they could not have any form of headache at the beginning of the experiment.

Migraine patients (n=32) were scanned: i) at baseline (outside an attack; ii) during randomly allocated and double-blind infusion of NTG (0.5 µg/kg/min over 20 min; n=27) or placebo (n=5); and iii) during an ensuing migraine attack or, if no migraine had occurred, at 6 hours after infusion. The duration of the scan sessions was approximately 25 minutes. The study subjects remained in the scanner between the baseline and the NTG or placebo infusion scanning sessions which began 10 minutes after onset of the infusion. Heart rate and blood pressure were monitored during the experiments. Two days after the experiment, subjects were contacted by telephone to check whether a migraine attack had occurred beyond the 6-hour time window ²²¹.

Placebo administration was included in the protocol to minimise patient and observer's bias for diagnosing whether or not NTG infusion had provoked a migraine headache (as this diagnosis is based on subjective assessment of symptoms ³). We choose for an unequal and incomplete allocation to receiving NTG or placebo mainly for two

regel 1 _____ reasons. First, NTG administration was only used as a well established tool to provoke
regel 2 _____ migraine attacks. Our study objective was primarily to assess intra-individual changes
regel 3 _____ from baseline, rather than comparing the effect of NTG with that of placebo. Secondly,
regel 4 _____ we wanted to minimise the number of patients who would contribute only very little to
regel 5 _____ the study results (placebo was only given for masking reasons) to reduce unnecessary
regel 6 _____ burden to patients, investigators, and MRI scanning time (the study protocol was very
regel 7 _____ time consuming).

regel 8 _____ ***Magnetic resonance angiography***

regel 9 _____ The MR investigations were performed on a 3.0-Tesla whole-body system (Philips
regel 10 _____ Medical Systems, The Netherlands). The MRA protocol consisted of two parts, one to
regel 11 _____ assess blood vessel diameter changes and one to assess blood flow changes.
regel 12 _____

regel 13 _____ The “blood vessel diameter protocol” consisted of a thick two-dimensional phase
regel 14 _____ contrast (2D PC) sagittal localiser survey through the circle of Willis, followed by a
regel 15 _____ three-dimensional time-of-flight (3D TOF) MRA sequence to visualise the BA and ECA,
regel 16 _____ ICA, PCA and MCA on both sides. This scan had the following imaging parameters:
regel 17 _____ repetition time / echo time (TR/TE): 22 ms / 3.5 ms; flip angle 15°; field of view: 220
regel 18 _____ x 220 mm; number of excitations: 1; slice orientation: transverse; slice thickness: 0.65
regel 19 _____ mm; number of slices: 200; scan percentage 100%, matrix reconstruction size: 512 x
regel 20 _____ 512 resulting in a nominal voxel size (x,y,z) of 0.43 x 0.43 x 0.65 mm; total acquisition
regel 21 _____ time: 4min 30sec. Based on the reconstruction of this 3D-TOF a second 3D-TOF with
regel 22 _____ a higher spatial resolution was performed to visualise the extra- and intracranial parts
regel 23 _____ of the MMA on both sides. This scan had the following imaging parameters: TR/TE: 15
regel 24 _____ ms / 2.1 ms; flip angle 15°; field of view: 200 x 200 mm; number of excitations: 1; slice
regel 25 _____ orientation: transverse; slice thickness: 0.25 mm; number of slices: 130; scan percentage
regel 26 _____ 100%, matrix reconstruction size: 512 x 512 resulting in a nominal voxel size (x,y,z) of
regel 27 _____ 0.39 x 0.39 x 0.25 mm; total acquisition time: 8min 31sec.
regel 28 _____

regel 29 _____ For the “blood flow protocol”, a 2-dimensional phase-contrast (2D-PC) section was
regel 30 _____ positioned on the basis of two thick slab localiser MRA scans in the coronal and sagittal
regel 31 _____ plane at the level of the skull base, perpendicular on the ICA and BA, to measure the
regel 32 _____ flow volume. The MRA flow volume measurements in the present study are derived
regel 33 _____ from previously developed and optimized protocols ²¹¹⁻²¹⁴. Acquisition parameters:
regel 34 _____ repetition time / echo time (TR/TE): 16 ms / 8.5 ms; flip angle 10°; field of view: 150
regel 35 _____ x 150 mm; number of excitations: 20; slice orientation: transverse; slice thickness: 5.0
regel 36 _____ mm; number of slices: 1; scan percentage 100%; PC velocity encoding: 140 cm/s; matrix
regel 37 _____
regel 38 _____
regel 39 _____

reconstruction size: 256 x 256 resulting in a nominal voxel size (x,y,z) of 0.59 x 0.59 x 50 mm; total acquisition time: 56sec. Figure 1 illustrates the positioning of the 2D PC section through the ICA and BA. On an independent workstation, quantitative flow values were calculated in each vessel by integrating across manually drawn regions of interest that enclosed the vessel lumen closely.

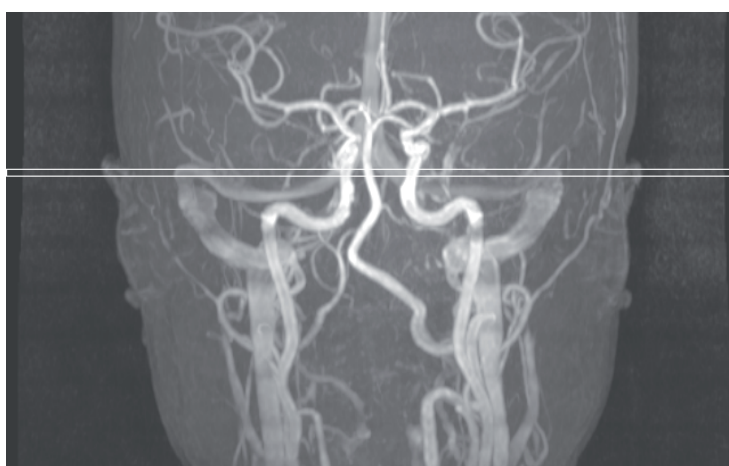


Figure 1 Magnetic resonance angiography, coronal maximum intensity projection. Horizontal line indicates the positioning of the 2-dimensional phase-contrast section through the ICA and the BA.

Image post processing: diameter calculations

All MRA images were transferred to a remote workstation for quantitative analysis using the Quantitative-MRA (QMRA) software package developed at our institution. A full description of the contour detection methods used and the validation have been described previously²¹⁵. The software provides automated contour detection and quantification of the luminal boundaries in selected vessel segments in 3D MRA datasets. The only user interaction required is the definition of the vessel segment of interest by placing a proximal and distal point in the 3D dataset. Subsequently, the software detects a 3D path line following the centre of the vessel lumen and cross-sectional multiplanar reconstructions (MPR's) are generated perpendicular to the centreline at 0.5 mm intervals. In each of these MPR's a contour around the vessel lumen is detected automatically. From these contours, based on the assumption of circular vessel cross-sections, the average diameter of the selected vessel segment is derived. Blood vessel segments were selected as follows: A) the MMA was measured in an extra-cranial segment (from the origin at the maxillary artery to the end, 5 to 6 mm distally; Figure2); B) the ECA from

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

the origin at the superficial temporal artery to the end, 10 mm proximally; C) the ICA from just proximally of the syphon to the end, 15 mm distally; D) the MCA, onset after A1 segment and end 8 mm distally; E) the BA, from the origin at the PCA to the end 12 mm proximally; F) the PCA, beginning at the origin at BA and end 8 mm distally). Location of measured vessel segments were kept constant within subjects.



Figure 2 Magnetic resonance angiography of the MMA region and position of the measured segment: (A) maxillary artery, (B) middle meningeal artery (MMA).

Statistical analysis

We first tested the left-to-right differences in diameters for bilateral blood vessels (MMA, ICA, ECA, MCA and PCA) using paired t-tests. Since the differences were not statistically significant, we only present the mean diameters for the right and left blood vessels throughout the manuscript. The effect of NTG and migraine attack on blood vessel diameters and blood flow were tested using a linear mixed model. Patients with a migraine attack (n=20) were compared to patients without an attack after NTG (n=7). Data from patients receiving placebo were not used for statistical testing. $P < 0.05$ was considered statistically significant.

RESULTS

Clinical effects of infusion of NTG or placebo

In total 32 migraine patients were randomly infused with either NTG (N=27) or placebo (N=5). Demographic characteristics of the study population are summarised in Table 1. No attack occurred after placebo (0/5). In contrast, infusion of NTG provoked a migraine attack (all without aura) in 20/27 (74%) migraine patients after a median time of 3.75 hours (range: 1.5 - 5.5 hours). In 18/20 attacks the headache was unilateral (left: n=9; right n=9). The clinical characteristics of the patients who developed a migraine attack in response to NTG and the clinical features of the provoked attacks are summarised in Supplemental Table s1.

Table 1 Demographic characteristics of study participants

Intervention	Migraine (n=32)		
	NTG (27)	Placebo (n=5)	
Attack	Yes (n=20)	No (n=7)	No
Age in years (SD)	45.5 (8.5)	34 (8.9)	44.8 (13.3)
Ratio female to males	15 : 5	7 : 0	3 : 2
Ratio MO to MA	17 : 3	6 : 1	4 : 1
Attack frequency; mean (SD)	2.6 (1.0)	2.1 (0.38)	2.4 (1.1)

MO denotes migraine without aura, MA migraine with aura.

Side to side differences for blood vessel diameters

There were no ($p>0.05$) right-to-left differences for the diameters of the four bilateral blood vessels (MMA, ICA, ECA, MCA, PCA) in any of the three conditions (data not shown), except for the MCA during session three ($p=0.024$). This difference was considered not significant after correction for multiple testing. Similarly, in the 18 patients with a unilateral headache, there were no significant ($p>0.05$) differences between the diameters on the headache and the non-headache side (Supplemental Table s4). Therefore, the mean diameters of the right and left blood vessels are presented throughout the paper.

Diameter and blood flow changes during infusion of NTG or placebo

During NTG infusion there was a significant vasodilatation of all blood vessels compared to baseline (Figures 3A to F and Supplemental Table s2; $p<0.01$ for all blood vessels). The diameter increase was greatest in the extra-cerebral blood vessels (MMA and ECA), ranging from 16.4% to 30.3%, as compared to 6.7% - 20.7% diameter increase in the

intra-cranial blood vessels (ICA, MCA, BA and PCA). During infusion of placebo, there were no changes in diameter for any of the blood vessels. There were no changes in ICA or BA blood flow during infusion of NTG or placebo (Figure 4A – B and Supplemental Table s3).

Figure 3A -F Mean blood vessel diameter changes (mean of left and right in bilateral vessels) in six selected intracranial blood vessels at baseline, during infusion of nitroglycerin (NTG) or placebo, and during an NTG-provoked migraine or, if no attack had occurred, at 6 hours after infusion. (● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack).

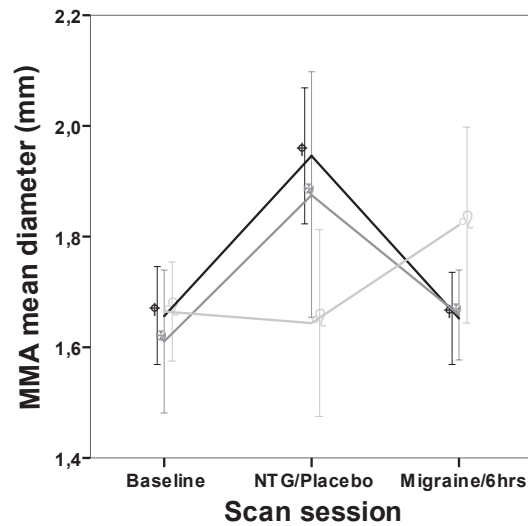


Figure 3A

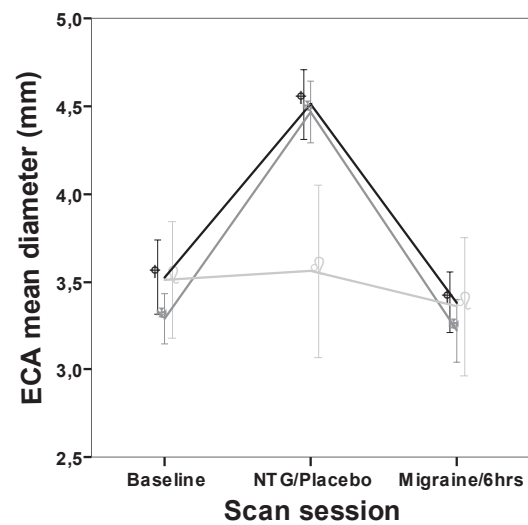


Figure 3B

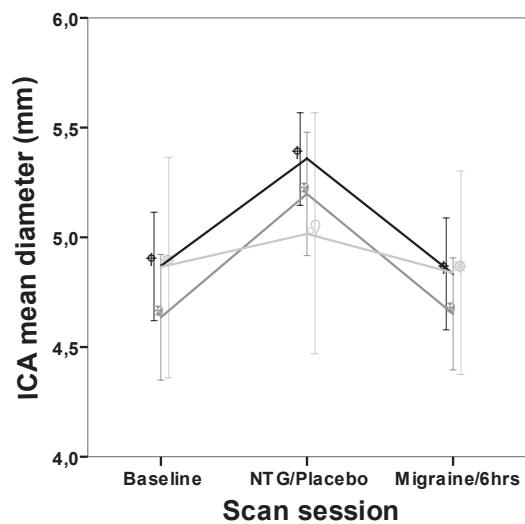


Figure 3C

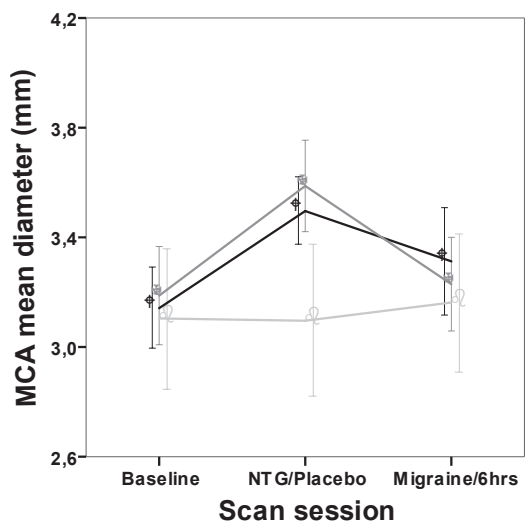


Figure 3D

- regel 1
- regel 2
- regel 3
- regel 4
- regel 5
- regel 6
- regel 7
- regel 8
- regel 9
- regel 10
- regel 11
- regel 12
- regel 13
- regel 14
- regel 15
- regel 16
- regel 17
- regel 18
- regel 19
- regel 20
- regel 21
- regel 22
- regel 23
- regel 24
- regel 25
- regel 26
- regel 27
- regel 28
- regel 29
- regel 30
- regel 31
- regel 32
- regel 33
- regel 34
- regel 35
- regel 36
- regel 37
- regel 38
- regel 39

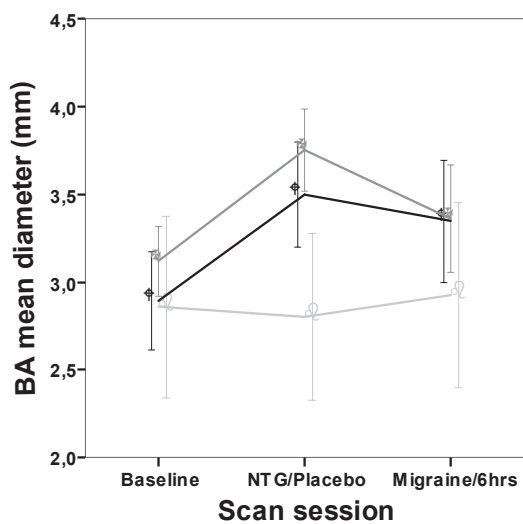


Figure 3E

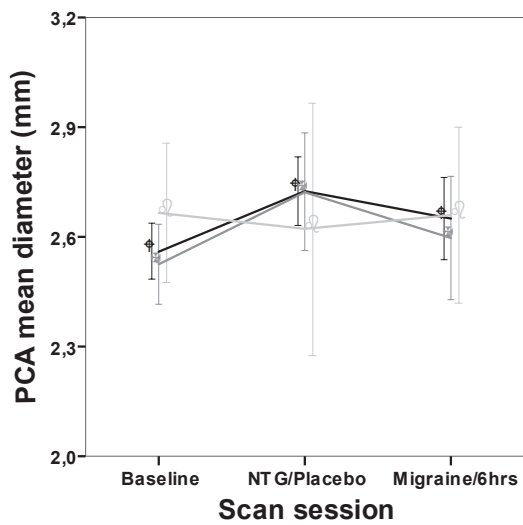


Figure 3F

Table 2 Mean blood vessel diameters (mean of right and left for bilateral blood vessels) of six selected intracranial blood vessel at baseline and during an NTG-provoked migraine attack or, if no attack had occurred, at 6 hours after infusion in 32 migraine patients.

Blood vessel	Inter-vention	Migraine attack	N	A)	B)	Change (B vs A) mm (% from A)
				Baseline mm (SD)	During migraine or at 6 hours mm (SD)	
MMA	NTG	Yes	20	1.66 (0.19)	1.65 (0.17)	-0.01 (-0.6)
	NTG	No	7	1.61 (0.12)	1.66 (0.08)	0.05 (3.1)
	Placebo	No	5	1.67 (0.73)	1.82 (0.14)	0.16 (9.6)
ECA	NTG	Yes	20	3.53 (0.42)	3.38 (0.36)	-0.12 (-3.4)
	NTG	No	7	3.29 (0.16)	3.22 (0.19)	-0.07 (-2.1)
	Placebo	No	5	3.51 (0.27)	3.36 (0.32)	-0.15 (-4.3)
ICA	NTG	Yes	20	4,87 (0.53)	4,83(0.53)	-0.04 (-0.8)
	NTG	No	7	4,64 (0.31)	4,65(0.28)	0.01 (0.2)
	Placebo	No	5	4,86 (0.41)	4,84(0.37)	-0.02 (-0.4)
MCA	NTG	Yes	20	3,14 (0.32)	3,31 (0.41)	0.17 (5.4)
	NTG	No	7	3,19 (0.19)	3,23 (0.19)	0.04 (1.3)
	Placebo	No	5	3,10 (0.20)	3,16 (0.20)	0.06 (1.9)
BA	NTG	Yes	20	2,89 (0.60)	3,35 (0.72)	0.48 (16.6)
	NTG	No	7	3,12 (0.21)	3,36 (0.33)	0.24 (7.7)
	Placebo	No	5	2,86 (0.42)	2,93 (0.42)	0.07 (2.5)
PCA	NTG	Yes	20	2.56 (0.16)	2.65 (0.23)	0.09 (3.5)
	NTG	No	7	2.52 (0.12)	2.60 (0.18)	0.07 (2.8)
	Placebo	No	5	2.67 (0.15)	2.66 (0.19)	0.01 (0.4)

NTG denotes nitroglycerin, MMA middle meningeal artery, ECA external carotid artery, ICA internal carotid artery, MCA middle cerebral artery, BA basilar artery and PCA posterior cerebral artery. There were no significant changes in diameter during the migraine attack.

Table 3 Blood flow in the basiliary (BA) and internal carotid artery (ICA) (mean of left and right) in migraine patients at baseline and during a migraine attack or, if no attack had occurred, at 6 hours after infusion of NTG or placebo.

Blood vessel	Inter-vention	Migraine Attack	N	A)	B)	Difference (B vs A)
				Blood flow Baseline	Blood flow During Migraine or at 6 hours	
				ml/min (SD)	ml/min (SD)	ml/min
BA	NTG	Yes	20	173.7 (69.4)	128.5 (40.1)	-46.2
	NTG	No	7	177.2 (71.9)	189.7 (26.3)	12.5
	Placebo	No	5	170.5 (39.4)	176.9 (63.6)	6.4
ICA	NTG	Yes	20	589.6 (128.5)	542.0 (166.8)	-57.7
	NTG	No	7	542.9 (101.2)	468.6 (151.2)	-74.3
	Placebo	No	5	542.0 (211.1)	522.8 (276.7)	-19.2
Total cerebral blood flow	NTG	Yes	20	763.3 (124.1)	670.5 (166.6)	-92.8
	NTG	No	7	720.1 (97.7)	658.3 (153.4)	-61.8
	Placebo	No	5	712.6 (202.4)	699.7 (253.9)	-12.8

NTG denotes nitroglycerin, BA basiliary artery and ICA internal carotid artery. Difference between patients with an attack compared to patients without an attack after NTG were not significant.

Diameter and blood flow changes during migraine attacks

Compared to baseline, there were no significant ($p > 0.05$) diameter changes during attacks for any of the blood vessels (Table 2 and Figures 3A to F). This was also true when controlling for the headache side in the 18 patients with an unilateral headache; the changes on the headache side were no different compared to those on the non-headache side (Supplemental Table s4). Similarly, there were no significant ($p > 0.05$) differences when comparing the mean diameter changes (baseline vs. attack) in the 20 patients who developed a migraine attack after NTG with the changes (baseline vs. 6 hours post infusion) in the 7 patients who did not develop an attack and were measured 6 hours after infusion. The attack vs. no-attack change-differences were for the MMA = 0.06 mm (95% CI: -0.8; 0.21), for the ECA = 0.05 mm (95% CI: -0.14; 0.24), for the ICA = 0.06 mm (95% CI: -0.19; 0.31), for the MCA = -0.13 (95% CI: -0.41; 0.14), for the BA = -0.24 (95% CI: -0.59; 0.11), and for the PCA = -0.02 (95% CI: -0.22; 0.18). There were also no significant ($p > 0.05$) changes in total-, BA-, or ICA-blood flow during a migraine

attack when compared to baseline, nor were there significant ($p>0.05$) differences in the changes observed during attacks when compared to the changes in the patients who did not develop an attack and were measured 6 hours after infusion (Supplemental Table 4).

Figure 4A -B Mean blood flow in ICA (mean of left and right) and BA at baseline, during infusion of nitroglycerin (NTG) or placebo, and during an NTG-provoked migraine or, if no attack had occurred, at 6 hours after infusion. (● Migraine patients (NTG) with a provoked attack, ▲ Migraine patients (NTG) without an attack, X Migraine patients (placebo) without an attack)

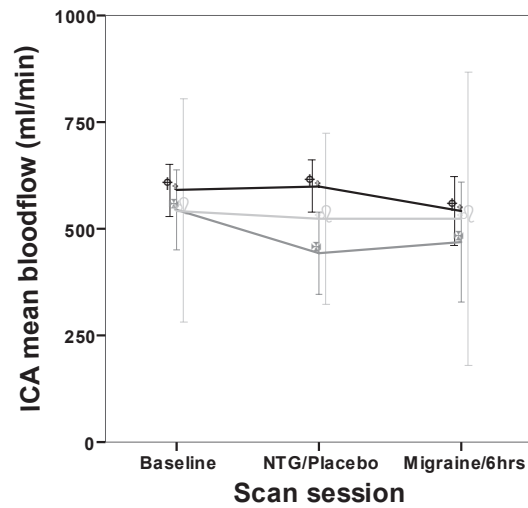


Figure 4A

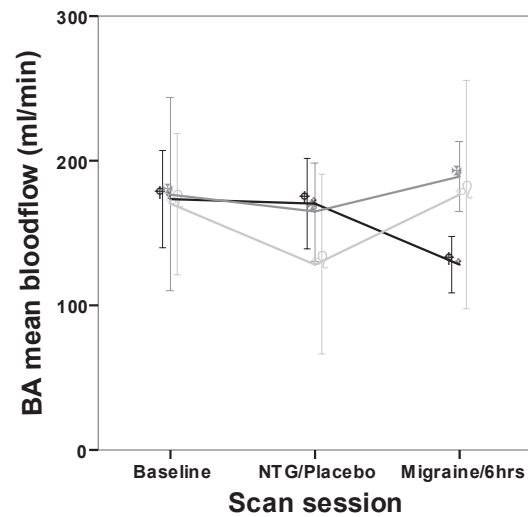


Figure 4B

regel 1
 regel 2
 regel 3
 regel 4
 regel 5
 regel 6
 regel 7
 regel 8
 regel 9
 regel 10
 regel 11
 regel 12
 regel 13
 regel 14
 regel 15
 regel 16
 regel 17
 regel 18
 regel 19
 regel 20
 regel 21
 regel 22
 regel 23
 regel 24
 regel 25
 regel 26
 regel 27
 regel 28
 regel 29
 regel 30
 regel 31
 regel 32
 regel 33
 regel 34
 regel 35
 regel 36
 regel 37
 regel 38
 regel 39

DISCUSSION

We used a well established NTG provocation model to induce faithfully migraine attacks and a highly sensitive, non-invasive 3T MRA technique to visualise and measure even small intra-individual diameter changes of cerebral and meningeal blood vessels. Contrary to longstanding and widespread belief, we failed to detect any evidence for a clinically relevant vasodilatation of major cerebral or meningeal blood vessels during migraine attacks. This finding has important implications for the understanding of the pathophysiology of the migraine headache and the development of future antimigraine agents. Novel antimigraine treatments may not require vasoconstrictor activity as predicted earlier ⁶¹.

In our provocation experiments, we infused NTG over a 20 min period and observed a vessel-dependent 7 – 30% vasodilatation at 10 minutes after beginning of the infusion. The vasodilatory effect is believed to be due to a direct local effect of NO on vascular smooth muscle cells ²⁴⁰ or through the release of vasoactive peptides such as CGRP ^{114,241}. Our findings on the early vascular effect of NTG are in accordance with those of ¹⁰⁹. Using 1.5T MRA they found a peak vasodilatation at 10 - 15 minutes after beginning of the NTG infusion and a normalisation of the vascular diameters back to baseline at 45 minutes after stopping of the infusion. For logistic reasons, we did not scan at 45 min after the infusion to confirm normalisation of the blood vessel diameter. However, in view of the well known short duration of action of NTG ²⁴² and the observed time course of the early vascular responses by ¹⁰⁹, we feel confident that blood vessel diameters had returned to baseline by one hour after the second (infusion) scan. It therefore seems justified to compare measurements during attacks with those obtained at baseline, before infusion.

The most important finding of the present study is that migraine headache was not associated with a clinically relevant vasodilatation of major cerebral or meningeal blood vessels, not even when controlled for headache side. We feel confident that this was not due to too low a sensitivity of the detection method. The very fact that we were able to detect an early transient vasodilatation in response to NTG of as low as 7% shows that the method we used is sufficiently sensitive to measure even small diameter changes. The clinical relevance of smaller changes is doubtful as during NTG infusion we observed an up to 30% increase in blood vessel diameter without associated migraine headache. Our results are also in agreement with at least some older TCD studies failing to show blood velocity changes indicative for vasodilatation during migraine attacks. ²⁴³⁻²⁴⁶

Finally, BA and ICA blood flow did also not change during migraine attacks. Cerebral blood flow is dependent on cardiac output, arterial caliber, and vasomotor tone in small resistance vessels.²⁴⁷ As blood pressure (as a measure for cardiac output; data not shown) and the BA and ICA diameters had not changed, it seems likely that there were also no changes in the intracranial resistance microvasculature during migraine attacks. In conclusion, our data seem to refute an important role of cerebral or meningeal vasodilatation in causing migraine headache. This would certainly be in accordance with observations that non-vascular mechanisms, such as exposure to sildenafil,⁵⁸ are capable of inducing migraine attacks.

Potential limitations of our study include that we didn't measure just before or at the onset of the migraine headache. We could thus have missed a brief transient vasodilatation at the very beginning of the migraine headache. Although unlikely, we cannot exclude this possibility. Another important question is whether and to what extent NTG-provoked migraine attacks are similar to spontaneous attacks. There are strong clinical and pathophysiological arguments in favour of this notion. The clinical symptoms and features, including the occurrence of premonitory symptoms several hours before the headache⁵⁶ and the response to anti-migraine drugs²³⁹, are strikingly similar between spontaneous and NTG-induced attacks. Likewise, in both there is an increase of CGRP in jugular venous blood^{61,124} and activation of the dorsal rostral brainstem on positron emission tomography.^{82,202} The fact that NTG provokes migraine aura's only rarely, even in patients with migraine with aura^{162 248}, seems to point at a trigger site of action beyond the aura triggering mechanism. We thus feel confident that our findings in NTG-provoked attacks can be extrapolated to spontaneous migraine headaches.

In this study, we did not observe significant changes in blood vessel diameter or blood flow during the headache phase of provoked migraine attacks. However, there were some (non-significant) changes in the posterior circulation that need to be discussed. First, the diameter of the BA did not return to baseline levels, unlike the other blood vessels. This was, however, true for both patients who had developed a delayed migraine headache and for those who had not. Secondly, the blood flow in the BA was decreased (although not significantly) from 174 ml/min at baseline to 129 ml/min in patients who had developed a migraine headache after GTN, whilst there was no such change in patients who had not developed a migraine headache. Whether these findings are clinically relevant, needs to be explored. A tentative correlation, for instance, could be made with previous findings of In previous studies our group has shown our group demonstrating increased prevalence of pontine hyperintensities and cerebellar infarcts in migraineurs from the general population^{249 250}.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

We conclude that, contrary to a longstanding and widespread belief, cerebral and meningeal diameter changes in migraine attacks, if at all happening, appear not to be of primary importance to the pathophysiology of the migraine headache.

SUPPLEMENTAL TABLES

Table s1 (only for publication on the web)
Characteristics of the NTG provoked migraine attack per subject

Subject	Sex	Age	Attack freq (per month)	Characteristics of provoked attack								Time to attack (hours)
				HS	UH	PH	AH	N	V	PT	PN	
1	M	40	2	2			+			+	+	4.5
2	M	35	3	2	+	+		+	+			2.5
3	F	42	2	2	+	+	+	+				4.5
4	M	54	4	2	+	+	+	+				4
5	F	49	4	2		+	+	+		+	+	4.5
6	F	55	2	2	+		+	+		+	+	3
7	F	48	1	2	+	+	+			+	+	3
8	F	37	2	2	+	+		+	+	+	+	2.5
9	F	33	2	2	+		+	+	+	+	+	4
10	F	32	2	2	+	+	+	+			+	5
11	F	51	3	2	+	+	+	+		+	+	3
12	F	28	3	2	+		+			+	+	3
13	F	55	4	2	+	+	+	+		+		5.5
14	F	55	4	2	+	+	+			+	+	2
15	F	53	2	2	+		+			+	+	5
16	M	46	3	2	+		+	+	+	+		2
17	M	51	4	2	+		+	+	+	+		4
18	F	49	0.5	2	+		+			+	+	4
19	F	50	4	2	+		+	+	+	+	+	3.5
20	F	31	1	2	+	+	+	+	+	+	+	1.5

F denotes female, M male, HS headache severity (2=moderate), UH unilateral headache (+ indicates yes, empty box no), PH pulsating headache, AH aggravation of headache during physical activity, N nausea, V vomiting, PT photophobia, PN phonophobia

Migraine headache is not associated with cerebral or meningeal vasodilatation

Table s2 (only for publication on the web)

Mean blood vessel diameters (mean of right and left for bilateral blood vessels) of six selected intracranial blood vessel at baseline and during infusion of nitroglycerin or placebo in 32 migraine patients.

Blood vessel	Inter-vention	N	A)	B)	Change (B vs A) mm (% from A)
			Baseline mm (SD)	During NTG or placebo mm (SD)	
MMA	NTG	27	1.65 (0.18)	1.93 (0.24)	0.27 (16.4)*
	Placebo	5	1.67 (0.07)	1.64 (0.12)	-0.02 (-1.2)
ECA	NTG	27	3.46 (0.38)	4.50 (0.38)	1.05 (30.3)*
	Placebo	5	3.51 (0.27)	3.56 (0.39)	0.05 (1.4)
ICA	NTG	27	4.81 (0.49)	5.32 (0.42)	0.51 (10.6)*
	Placebo	5	4,86 (0.41)	5,02 (0.44)	0.15 (3.1)
MCA	NTG	27	3.16 (0.29)	3.52 (0.24)	0.37 (11.7)*
	Placebo	5	3,10 (0.20)	3,10 (0.22)	-0.01 (-0.3)
BA	NTG	27	2.95 (0.53)	3.56 (0.57)	0.61 (20.7)*
	Placebo	5	2,86 (0.42)	2,80 (0.38)	-0.06 (-2.1)
PCA	NTG	27	2.55 (0.15)	2.72 (0.19)	0.17 (6.7)*
	Placebo	5	2.67 (0.15)	2.62 (0.28)	-0.04 (-1.5)

NTG denotes nitroglycerin, MMA middle meningeal artery, ECA external carotid artery, ICA internal carotid artery, MCA middle cerebral artery, BA basilar artery, PCA posterior cerebral artery. * NTG effect on diameter was significant in all six blood vessels (p<0.01).

Table s3 (only for publication on the web)

Blood flow in BA, ICA and total cerebral blood flow in migraine patients at baseline and during infusion of nitroglycerin or placebo.

Blood vessel	Inter-vention	N	A)	B)	Change (B vs A)
			Baseline	During NTG or placebo	
			ml/min (SD)	ml/min (SD)	ml/min (%)
BA	NTG	27	174.6 (68.7)	169.3 (57.9)	-5.4 (-3.1)
	Placebo	5	170.5 (39.4)	128.6 (49.9)	-41.9 (-24.6)
ICA	NTG	27	577.1 (121.6)	557.5 (139.4)	-19.5 (-3.4)
	Placebo	5	542.0 (211.1)	523.2 (161.9)	-18.8 (-3.5)
TCBF	NTG	27	751.7 (117.3)	726.8 (149.5)	-24.9 (-3.3)
	Placebo	5	712.6 (202.4)	651.8 (198.8)	-60.7 (-8.5)

NTG nitroglycerin, ICA internal carotid artery, BA basilar artery, tCBF total cerebral blood flow.

Table s4 (only for publication on the web)

Blood vessel diameter of five bilateral intracranial blood vessels at baseline and during an NTG-provoked migraine attack in 18 migraine patients with unilateral headache.

Blood vessel	Side	A)	B)	Change (B vs A)
		Baseline	During migraine	
		mm (SD)	mm (SD)	mm (% from A)
MMA	Headache	1.69 (0.22)	1.67 (0.21)	-0.03 (-1.78)
	Non-headache	1.60 (0.18)	1.58 (0.17)	-0.03 (-1.88)
ECA	Headache	3.51 (0.39)	3.34 (0.38)	-0.18 (-5.13)
	Non-headache	3.43 (0.46)	3.33 (0.39)	-0.04 (-1.17)
ICA	Headache	4.87 (0.59)	4.79 (0.64)	-0.09 (-1.85)
	Non-headache	4.89 (0.55)	4.87 (0.55)	-0.02 (-0.41)
MCA	Headache	3.19 (0.34)	3.32 (0.44)	0.13 (4.08)
	Non-headache	3.15 (0.34)	3.36 (0.45)	0.24 (7.62)
PCA	Headache	2.58 (0.19)	2.72 (0.30)	0.13 (5.0)
	Non-headache	2.58 (0.21)	2.66 (0.23)	0.08 (3.1)

MMA denotes middle meningeal artery, ECA external carotid artery, ICA internal carotid artery, MCA middle cerebral artery and PCA posterior cerebral artery. There were no significant differences in diameter change between headache side and non-headache side.

GENERAL DISCUSSION AND CONCLUSIONS



- regel 1
- regel 2
- regel 3
- regel 4
- regel 5
- regel 6
- regel 7
- regel 8
- regel 9
- regel 10
- regel 11
- regel 12
- regel 13
- regel 14
- regel 15
- regel 16
- regel 17
- regel 18
- regel 19
- regel 20
- regel 21
- regel 22
- regel 23
- regel 24
- regel 25
- regel 26
- regel 27
- regel 28
- regel 29
- regel 30
- regel 31
- regel 32
- regel 33
- regel 34
- regel 35
- regel 36
- regel 37
- regel 38
- regel 39

regel 1 _____ This thesis deals with the association between potential trigger factors and the occurrence
regel 2 _____ of a migraine attack as well as the action mechanism of trigger factors in migraine. We
regel 3 _____ have focused our research on three trigger factors; mental stress, normobaric hypoxia
regel 4 _____ and nitroglycerin. The most remarkable findings will be summarized and discussed.

regel 5 _____
regel 6 _____ ***Premonitory symptoms are frequently reported by migraine patients in***
regel 7 _____ ***a clinic based sample (chapter 1)***

regel 8 _____ In a clinic based sample of 374 migraine patients we found that 86.9% of patients
regel 9 _____ reported at least one symptom and 71.1% reported two or more. Forty-nine patients
regel 10 _____ (13.1%) reported no premonitory symptoms. Other clinic based studies found prevalence
regel 11 _____ rates for premonitory symptoms of 79%¹³³ and 84%⁷, although rates of around 33%
regel 12 _____ also have been found^{134,135}. The most frequently reported premonitory symptoms were
regel 13 _____ fatigue (46.5%), phonophobia (36.4%) and yawning (35.8%). This is in accordance
regel 14 _____ with other studies assessing premonitory symptoms. In a prospective study in 100
regel 15 _____ migraine patients the most frequent symptoms were anxiety, phonophobia, irritability,
regel 16 _____ unhappiness and yawning⁷. Several important questions regarding premonitory symptoms
regel 17 _____ in migraine remain to be answered. For instance how specific are premonitory symptoms
regel 18 _____ for migraine? Many possible premonitory symptoms are rather a-specific and are also
regel 19 _____ associated with other conditions such as premenstrual syndrome¹³⁶ and depression¹⁴⁰.
regel 20 _____ Future studies could try to assess the occurrence of possible premonitory symptoms in
regel 21 _____ combination with other migraine symptoms in a prospective design using electronic
regel 22 _____ diaries in an unselected migraine population¹³¹.

regel 23 _____
regel 24 _____ ***The association between mental stress and the occurrence of a migraine***
regel 25 _____ ***attack is less clear than previously assumed (chapter 2)***

regel 26 _____ In chapter 2 we described findings of a prospective longitudinal study. We failed to
regel 27 _____ find any objective evidence for a temporal relationship between changes in perceived
regel 28 _____ stress, biological indicators for a stress-response, and the onset of a migraine attack.
regel 29 _____ Although stress-sensitive patients indeed reported an increase in perceived stress in
regel 30 _____ the days before an attack, this was not accompanied by objective signs indicating
regel 31 _____ a biological stress response. These findings are in accordance with previous stress
regel 32 _____ studies. Retrospective and prospective questionnaire studies suggest an increase in the
regel 33 _____ perception of potential stressors around the migraine attack^{27,42}. On the other hand
regel 34 _____ stress provocation studies including biological stress response measures during and
regel 35 _____ outside a migraine attack are either negative or not conclusive¹⁴¹. Furthermore, there is
regel 36 _____ also no association between migraine and major life stressors²⁵¹. Possibly mental stress
regel 37 _____ is not a trigger factor for migraine but do migraine patients perceive daily hassles as
regel 38 _____

stressful due to the development of a migraine attack. The lack of a biological stress response (eg cortisol) in patients who perceived more stress could be the result of the temporal resolution of our measurements as discussed in chapter 2. Future studies will have to answer the question whether mental stress is a trigger factor for migraine or not. An option could be to design an experimental provocation study using a mental stressor sufficient enough to trigger migraine attacks.

Normobaric hypoxia is a possible trigger factor for migraine (chapter 3)

A high altitude environment causes acute mountain sickness in healthy volunteers⁹⁷ and might be able to trigger migraine in susceptible patients¹²⁶, mainly through hypoxia, although an additional effect of hypobaria can not be excluded¹⁷⁹. Several symptoms in acute mountain sickness are comparable to migraine including headache, nausea, fluid retention and disturbance of sleep⁹⁷ and as we have shown in chapter 3 normobaric hypoxia triggered a migraine attack in 6 out of 14 migraine patients. The results for hypoxia were not significant but in the light of a very low response after the positive control nitroglycerin it could be that the studied population was not very susceptible for migraine. Combined with previous studies in mountaineers⁹⁹ it was concluded that hypoxia is a possible trigger factor for migraine. For the dutch migraine population there are no direct consequences of this finding, since the highest mountain in the Netherlands is only 321 meters. Airflights might cause a problem; current regulations require that cabin air pressure must be no lower than the air pressure that naturally occurs at 2400 meter. In our experiments subjects were exposed to hypoxia corresponding to 4500m. Whether 2400m would be enough to trigger migraine is unclear.

The migraine response after NTG infusion is variable ranging between 20% and 83% (chapter 3 and 6)

The migraine response after infusion of a nitric oxide donor ranges from 20% to 83% between studies. The reason for the observed variability is unclear. A low baseline attack frequency could be a likely explanation. A study comparing the migraine response in patients with <4 attacks per year with patients 12 attack per year showed a trend towards more migraine attacks in frequent sufferers.²⁵² The second explanation could be the occurrence of aura's. Two studies found a lower migraine response in patients with aura; 67% MA vs 83% MO⁵⁶ and 31.8% MA vs 78% MO⁵⁵. A third factor could be age. In the three studies with the lowest migraine response the mean age is rather low ranging from 29.1¹²⁶ to 34.3 years.^{123,129} Also in our second NTG study in migraine patients the mean age in non-responders was 34 year whereas the mean age in responders was 45.5 year (chapter 6). Whether there are other factors involved in migraine susceptibility for NTG is unclear.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____ ***Normobaric hypoxia causes cerebral edema in healthy volunteers***
regel 2 _____ ***(chapter 4)***

regel 3 _____ Research in acute mountain sickness has mainly focused on the development of cerebral
regel 4 _____ edema. Severe AMS is associated with high altitude cerebral edema¹⁰⁴; whether edema
regel 5 _____ also occurs in mild cases of AMS was unclear. In chapter 4 we have shown that hypoxia
regel 6 _____ caused vasogenic edema in healthy volunteers irrespective of AMS symptoms and that
regel 7 _____ cytotoxic edema might be associated with severe AMS. Our findings are in accordance
regel 8 _____ with other studies which have been published while our study was drafted¹⁶⁷. These
regel 9 _____ findings in AMS might have implications for migraine. Approximately 30% of healthy
regel 10 _____ climbers develop AMS at 3000m altitude⁹⁷. Whether migraine patients are more
regel 11 _____ susceptible for AMS is unclear. One of the mechanisms involved in the development of
regel 12 _____ vasogenic and/or cytotoxic edema is Na(+)-K(+)-ATPase¹⁹⁰. A mutation in the Na(+)-K(+)-
regel 13 _____ ATPase gene was found in familial hemiplegic migraine²⁵³, possibly causing blood brain
regel 14 _____ barrier disruption and cerebral edema²⁵⁴. Future studies in migraine using hypoxia as a
regel 15 _____ trigger will have to show whether cerebral edema is involved in the pathophysiology of
regel 16 _____ the common types of migraine.

regel 17 _____ ***Nitroglycerin induced vasodilatation in both healthy volunteers and***
regel 18 _____ ***migraine patients (chapter 5 and 7) can be measured reliably with MRA***

regel 19 _____ Nitroglycerin is an exogenous donor of nitric oxide²⁵⁵ and causes vasodilatation either
regel 20 _____ through relaxation of vascular smooth muscles or through the release of CGRP¹¹⁴. A
regel 21 _____ common technique to measure the diameter of cerebral blood vessels in vivo in humans
regel 22 _____ is using trans-cranial Doppler. The advantages of TCD are that it is a continuous, cheap
regel 23 _____ and non-invasive measurement, however, the outcome is operator dependent and
regel 24 _____ TCD is an indirect measurement of the blood vessel diameter since it measures blood
regel 25 _____ flow velocity²³⁷. To study blood flow diameters in migraine patients the aim was to
regel 26 _____ have a direct and non-invasive method which would enable us to study blood vessel
regel 27 _____ as small as the middle meningeal artery. Using magnetic resonance we have been able
regel 28 _____ to reliably measure blood vessel diameters as shown in chapter 5 and 6. MRA images
regel 29 _____ were measured by two independent observers and the agreement between observers
regel 30 _____ was 0.74 (an intra class correlation of 0.7 or more is considered acceptable).The MMA
regel 31 _____ diameter increase during NTG in healthy volunteers in chapter 5 was 20.1%, whereas the
regel 32 _____ diameter increase in an other sample of healthy volunteers in chapter 6 was 19.9%.

The blood flow response but not vasodilatory response to nitroglycerin in migraine is related to the development of a delayed migraine attack (chapter 6)

Nitroglycerin caused a decrease in ICA blood flow without affecting BA blood flow. The ICA blood flow decreased significantly more in healthy volunteers as compared to migraine patients. Previous NTG studies in migraine showed either no difference in blood velocity decrease in the middle cerebral artery between migraine and controls¹¹⁸, or a more pronounced decrease in migraine patients¹¹⁷. An explanation for the difference in results could be that these studies did not take into account the occurrence of a provoked attack. As shown in this study the patients without an attack (several hours after NTG infusion) showed a decrease in ICA blood flow similar to healthy volunteers whereas in patients with an attack ICA blood flow did not decrease. Blood flow in the ICA is affected by several parameters; i) ICA blood vessel diameter, ii) cardiac output and iii) vasomotor tone in small resistance vessels. The ICA diameter increased during NTG infusion, but there was no difference between groups. Nitroglycerin has shown to decrease cardiac output²¹⁶. In this study we did not measure cardiac output but we did not observe a difference in blood pressure response during NTG infusion between groups (data not shown), suggesting that there was no difference in decrease of cardiac output between groups. So a difference in vasomotor tone of small resistance vessels might be the main explanation for the observed difference between patients with and without an attack.

There is no vasodilatation during the headache phase of a nitroglycerin provoked migraine attack (chapter 7)

This is the most important conclusion in this thesis. For many years there has been debate concerning vasodilatation in meningeal and cerebral arteries during the headache phase of a migraine attack. Studies by Wolff et al. showed that stimulation of cerebral and meningeal arteries caused headache and it was suggested that vasodilatation of cranial blood vessels was the cause for headache during a migraine attack¹⁹⁸. Vasoactive substances such as nitroglycerin can trigger migraine in susceptible patients⁷⁸ and triptans might exert their anti-migraine effect through vasoconstriction of cranial blood vessels⁷⁵. However, in vivo measurements in humans using transcranial Doppler (TCD) are not conclusive^{204,226,227,243-246}. In chapter 7 we have shown that there is no vasodilatation or change in cerebral blood flow during the headache phase of a provoked migraine attack as a model for spontaneous attacks. This finding does imply that future anti-migraine drugs do not have to constrict cerebral or meningeal blood vessels to treat the headache during a migraine attack.

— regel 1
 — regel 2
 — regel 3
 — regel 4
 — regel 5
 — regel 6
 — regel 7
 — regel 8
 — regel 9
 — regel 10
 — regel 11
 — regel 12
 — regel 13
 — regel 14
 — regel 15
 — regel 16
 — regel 17
 — regel 18
 — regel 19
 — regel 20
 — regel 21
 — regel 22
 — regel 23
 — regel 24
 — regel 25
 — regel 26
 — regel 27
 — regel 28
 — regel 29
 — regel 30
 — regel 31
 — regel 32
 — regel 33
 — regel 34
 — regel 35
 — regel 36
 — regel 37
 — regel 38
 — regel 39

CONCLUSIONS AND FUTURE PERSPECTIVES

Based on the studies presented in this thesis several conclusions can be drawn. The most important conclusion is that there is no vasodilatation of cranial arteries during the headache phase of a migraine attack. Future drug development research should focus on non-vascular structures to treat migraine headache. The second conclusion is that there is no clear association between mental stress and the occurrence of a migraine attack in spite of previous reports. Based on this study it does not make sense to advise migraine patients to avoid potential mental stressors as part of their therapeutic plan. The discrepancy between objective and subjective stress measures needs further study. The third conclusion is that hypoxia might trigger migraine in susceptible patients through a process which may involve development of cerebral edema. This conclusion is rather speculative and needs more study in migraine patients using hypoxia as an experimental trigger factor.

SAMENVATTING EN CONCLUSIES

— regel 1

— regel 2

— regel 3

— regel 4

— regel 5

— regel 6

— regel 7

— regel 8

— regel 9

— regel 10

— regel 11

— regel 12

— regel 13

— regel 14

— regel 15

— regel 16

— regel 17

— regel 18

— regel 19

— regel 20

— regel 21

— regel 22

— regel 23

— regel 24

— regel 25

— regel 26

— regel 27

— regel 28

— regel 29

— regel 30

— regel 31

— regel 32

— regel 33

— regel 34

— regel 35

— regel 36

— regel 37

— regel 38

— regel 39

regel 1 _____ In dit proefschrift wordt de relatie tussen mogelijke uitlokkende factoren en een migraine
regel 2 _____ aanval beschreven als mede het werkingsmechanisme van uitlokkende factoren. Er is met
regel 3 _____ name gekeken naar drie factoren: mentale stress, normobare hypoxie en nitroglycerine.
regel 4 _____ De belangrijkste bevindingen zullen hierna samengevat en bediscussieerd worden.
regel 5 _____

Migraine (introductie)

regel 6 _____ In het eerste deel van dit proefschrift worden de klinische verschijnselen van migraine
regel 7 _____ besproken evenals het mechanisme onderliggend aan een migraine aanval. Migraine is
regel 8 _____ een neurologische aandoening waarbij hoofdpijn in aanvallen optreedt. Deze aanvallen
regel 9 _____ duren 4 uur tot 3 dagen. De hoofdpijn is vaak eenzijdig, bonzend en neemt toe bij
regel 10 _____ bewegen. Tevens treedt er misselijkheid, overgeven en overgevoeligheid voor licht,
regel 11 _____ geluid en geuren op. Voorafgaand aan de hoofdpijnfase noemen veel patiënten het
regel 12 _____ optreden van zogenaamde prodromale verschijnselen, zoals concentratie problemen,
regel 13 _____ vocht vasthouden en stemmingsproblemen.
regel 14 _____

regel 15 _____ De aanvalsfrequentie kan uiteenlopen van 1 migraineaanval per jaar tot meerdere
regel 16 _____ aanvallen per maand. Ondanks vele jaren van onderzoek weet men niet waardoor
regel 17 _____ migraine aanvallen ontstaan. In de literatuur worden veel mogelijke uitlokkende
regel 18 _____ factoren genoemd, zoals verschillende voedingsproducten, veranderingen in het weer,
regel 19 _____ stress verhogende situaties en vrouwelijke hormonen. Of er inderdaad een causaal
regel 20 _____ verband is tussen de mogelijke uitlokkende factoren en het optreden van een migraine
regel 21 _____ aanval is onduidelijk. Tevens is het mechanisme leidend tot een migraine aanval voor
regel 22 _____ een groot deel onbekend. Tijdens de hoofdpijnfase van een migraineaanval raakt de
regel 23 _____ vijfde hersenzenuw geactiveerd, maar wat hieraan voorafgaat, is niet duidelijk. Mogelijk
regel 24 _____ speelt vaatverwijding van hersenbloedvaten een rol. Een ander mogelijk mechanisme
regel 25 _____ is een tijdelijk defect in de bloed-hersen barrière waardoor uitlopers van de vijfde
regel 26 _____ hersenzenuw geprikkeld worden.
regel 27 _____

Prodromale verschijnselen worden frequent gemeld door migraine patiënten (hoofdstuk 1)

regel 28 _____ In een populatie van 389 migraine patiënten is gekeken naar het voorkomen van
regel 29 _____ prodromale verschijnselen. Dit zijn verschijnselen die optreden voorafgaand aan de
regel 30 _____ hoofdpijnfase van een migraine aanval. De meest genoemde verschijnselen waren
regel 31 _____ vermoeidheid (46.5%), lichtschuwheid (36.4%) en gapen (35.8%). Het bleek dat 86.9%
regel 32 _____ van de patiënten tenminste 1 prodromaal verschijnsel noemden en 71.1% noemde
regel 33 _____ er twee of meer. De bevindingen komen overeen met resultaten uit eerdere studies.
regel 34 _____ Enkele belangrijke vragen blijven echter onopgelost. Het is bijvoorbeeld onduidelijk hoe
regel 35 _____ specifiek prodromale verschijnselen zijn voor het optreden van een migraine aanval. Er
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

bestaat bijvoorbeeld een aanzienlijke overlap tussen prodromale verschijnselen en het premenstrueel syndroom en depressiviteit. Goede prospectieve studies zijn nodig om de sensitiviteit en specificiteit van prodromale verschijnselen voor een migraine aanval te bepalen.

De relatie tussen mentale stress en het optreden van een migraine aanval is minder duidelijk dan voorheen aangenomen (hoofdstuk 2)

In hoofdstuk 2 worden de bevindingen van een prospectieve longitudinale studie naar de relatie tussen mentale stress en migraine beschreven. Ondanks aanwijzingen in de literatuur dat er een duidelijke relatie bestaat tussen stress en migraine liet deze studie geen duidelijk verband zien tussen veranderingen in subjectieve en objectieve stress parameters en het optreden van een migraine aanval. In een subgroep van subjectief stress gevoelige patiënten was er wel een relatie tussen waargenomen stress en het optreden van een migraine aanval, maar dit ging niet gepaard met veranderingen in objectieve stress maten, zoals cortisol. Wellicht is het zo dat er wel een relatie is tussen waargenomen stress en migraine, maar dat stress de aanval niet uitlokt. Het zou kunnen zijn dat migraine patiënten vlak voor een aanval gevoelig zijn voor stress omdat ze in de aanloop van een aanval zitten.

Normobare hypoxie is een mogelijke uitlokkende factor voor migraine (hoofdstuk 3)

Verblijf op grote hoogte in de bergen kan leiden tot hoogteziekte in gezonde vrijwilligers en kan eventueel een migraine aanval uitlokken in migraine gevoelige patiënten. Het meest belangrijke mechanisme is de hypoxie (te weinig zuurstof). Verschillende hoogteziekte symptomen kunnen ook tijdens een migraineaanval optreden zoals hoofdpijn, misselijkheid en slaapproblemen. In hoofdstuk 3 is beschreven hoe blootstelling aan hypoxie (vergelijkbaar met een hoogte van 4500m) gedurende 5 uur een migraine aanval provoceerde in 6 van de 14 migraine patiënten. Hoewel het resultaat niet significant is, lijkt het erop dat hypoxie een mogelijke uitlokkende factor is voor migraine. In Nederland is hypoxie geen belangrijke factor omdat we geen bergen van betekenis hebben, maar hypoxie tijdens een vliegreis zou een migraine aanval kunnen uitlokken. Officiële regels stellen dat de luchtdruk aan boord van een vliegtuig minimaal vergelijkbaar moet zijn met 2400 meter.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____ ***Normobare hypoxie veroorzaakt cerebraal oedeem in gezonde***
regel 2 _____ ***vrijwilligers (hoofdstuk 4)***

regel 3 _____ Onderzoek naar de pathofysiologie van hoogteziekte laat zien dat er tijdens ernstige
regel 4 _____ hoogteziekte cerebraal oedeem ontstaat. Of er tijdens milde hoogteziekte ook oedeem
regel 5 _____ ontstaat niet duidelijk. Om meer over de effecten van hypoxie op de hersenen te weten
regel 6 _____ te komen is een groep studenten bloot gesteld aan een experimenteel model voor
regel 7 _____ hoogteziekte waarbij ze gedurende 6 uur hypoxisch gemaakt werden. De hypoxie
regel 8 _____ tijdens het experiment was vergelijkbaar met een hoogte van 4500m. Zoals beschreven
regel 9 _____ in hoofdstuk vier blijkt er zogenaamd vasogeen oedeem (vocht rondom de cellen) op
regel 10 _____ te treden na hypoxie; onafhankelijk van het optreden van hoogteziekte symptomen.
regel 11 _____ Voorts treedt er cytotoxisch oedeem (vocht in de cellen) op in de groep met de meeste
regel 12 _____ klachten. De bevindingen in de gezonde vrijwilligers hebben mogelijk implicaties voor
regel 13 _____ migraine. Een van de mechanismen betrokken bij de ontwikkeling van cerebraal oedeem
regel 14 _____ is het Na-K-ATP ase dat weer een rol speelt in familiere hemiplegische migraine (type
regel 15 _____ 2).

regel 16 _____ ***De kans op het krijgen van een migraine aanval na toediening van***
regel 17 _____ ***nitroglycerine ligt tussen 20% en 83% (hoofdstuk 3 en 6)***

regel 18 _____ Nitroglycerine is een bekende uitlokkende factor voor migraine. In twee verschillende
regel 19 _____ studies is gebruik gemaakt van nitroglycerine (NTG) voor het uitlokken van migraine
regel 20 _____ aanvallen (hoofdstuk 3 en 6). In de eerste studie was het percentage patiënten dat
regel 21 _____ een aanval kreeg na NTG 20% en in de tweede studie was dit 74%. In de literatuur
regel 22 _____ zijn er zelfs percentages tot 83% beschreven. De oorzaak voor deze variatie in het
regel 23 _____ effect van nitroglycerine is niet goed te geven. Een eerste mogelijke verklaring zou een
regel 24 _____ lage basale aanvalsfrequentie kunnen zijn. In een Deense studie is het effect van NTG
regel 25 _____ vergeleken tussen patiënten met een lage aanvalsfrequentie (minder dan 4 aanvallen
regel 26 _____ per jaar) en een hoge aanvalsfrequentie (meer dan 12 aanvallen per jaar). Deze studie
regel 27 _____ liet een trend zien in de richting van meer aanvallen in de groep met veel aanvallen.
regel 28 _____ Een tweede verklaring zou het wel of niet optreden van visuele aura's kunnen zijn. Er
regel 29 _____ zijn verschillende studies waarin de kans op een migraine aanval na NTG kleiner is in
regel 30 _____ patiënten die last van hebben van aura's. Een derde factor zou leeftijd kunnen zijn. In de
regel 31 _____ tweede studie is de leeftijd in de groep die geen aanval krijgt gemiddeld 34 jaar, terwijl
regel 32 _____ de leeftijd in de groep waarin wel een aanval optreedt gemiddeld 45.5 jaar is.
regel 33 _____
regel 34 _____
regel 35 _____
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

Nitroglycerine geïnduceerde vaatverwijding kan betrouwbaar gemeten worden met behulp van magnetic resonance angiografie (MRA) in zowel gezonde vrijwilligers als migraine patiënten (hoofdstuk 5 en 7)

Nitroglycerine veroorzaakt vaatverwijding in zowel veneuze als arteriële bloedvaten. Een veelgebruikte techniek om vaatverwijding in het hoofd te meten is zogenaamde transcraniële Doppler (TCD) waarbij er met geluidsgolven informatie over de bloedstroom in een bloedvat verkregen wordt. TCD is een goedkope en niet invasieve methode, echter de uitkomst van de meting is afhankelijk van de persoon die de meting doet. In verband met deze beperkingen hebben we gekozen om bloedvatdiameters te meten door middel van MRA. Alvorens het onderzoek in migraine patiënten uit te voeren is bij gezonde vrijwilligers naar de betrouwbaarheid van de MRA meting gekeken. Zoals besproken in hoofdstuk 5 is er een hoge correlatie (0.74) tussen twee onafhankelijke waarnemers, zodat de conclusie getrokken kan worden dat MRA een betrouwbare methode is.

De verandering in bloedstroom, in tegenstelling tot de verandering in bloedvat diameter, tijdens toediening van nitroglycerine is geassocieerd met het optreden van een migraine aanval (hoofdstuk 6)

In hoofdstuk zes is gekeken naar het effect van nitroglycerine op cerebrale bloedvaten (zowel diameter als bloedstroom) in gezonde vrijwilligers en in migraine patiënten. Het bleek dat nitroglycerine een forse vaatverwijding in alle gemeten bloedvaten gaf. Er was geen verschil in vaatverwijdend effect tussen migraine patiënten en vrijwilligers. Daarnaast daalde de bloedstroom in de arteria carotis interna (ICA) en bleef de bloedstroom in de arteria basilaris (BA) gelijk. In gezonde vrijwilligers daalde de bloedstroom veel sterker dan in migraine patiënten. Deze bevinding is in contrast met eerdere studies waarin de bloedstroom niet veranderde of juist meer veranderde in migraine patiënten. Een oorzaak voor het verschil zou kunnen zijn dat in eerdere studies geen onderscheid is gemaakt tussen patiënten met of zonder migraine aanval volgend op de NTG provocatie. Het bleek namelijk dat de ICA in patiënten zonder een migraine aanval sterk daalde (vergelijkbaar met gezonde vrijwilligers), terwijl de ICA bloedstroom in patiënten met een aanval zelf mild toenam. Bloedstroom in de ICA wordt bepaald door de ICA diameter, cardiac output en vasomotor tone in kleine weerstandsvaten. De diameter van de ICA nam toe, maar er was geen verschil tussen patiënten met of zonder aanval. Van nitroglycerine is bekend dat het hartminuut volume daalt kort na toediening, maar in deze studie is geen verschil gevonden in verandering in bloeddruk tussen patiënten met en zonder een migraine aanval. Dan blijft een mogelijk verschil in vasomotor tone in kleine weerstandsvaten over als verklaring voor het verschil.

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

regel 1 _____
regel 2 _____
regel 3 _____
regel 4 _____
regel 5 _____
regel 6 _____
regel 7 _____
regel 8 _____
regel 9 _____
regel 10 _____
regel 11 _____
regel 12 _____
regel 13 _____
regel 14 _____
regel 15 _____
regel 16 _____
regel 17 _____
regel 18 _____
regel 19 _____
regel 20 _____
regel 21 _____
regel 22 _____
regel 23 _____
regel 24 _____
regel 25 _____
regel 26 _____
regel 27 _____
regel 28 _____
regel 29 _____
regel 30 _____
regel 31 _____
regel 32 _____
regel 33 _____
regel 34 _____
regel 35 _____
regel 36 _____
regel 37 _____
regel 38 _____
regel 39 _____

Er treedt geen vaatverwijding op in cerebrale bloedvaten tijdens de hoofdpijnfase van een door nitroglycerine geïnduceerde migraine aanval (hoofdstuk 7)

Er wordt reeds vele jaren gediscussieerd over de relatie tussen vaatverwijding in hersenvaten en migraine. Wolff et al. liet zien dat stimulatie van bloedvaten in de hersenen en hersenvliezen erg pijngevoelig zijn. Voorts kunnen migraine aanvallen uitgelokt worden door vasoactieve middelen zoals nitroglycerine. Echter, studies met transcraniële Doppler, om vaatverwijding tijdens de hoofdpijn fase aan te tonen zijn niet eenduidig. De conclusie van hoofdstuk zeven is dat tijdens de hoofdpijnfase van een door nitroglycerine uitgelokte migraineaanval geen vaatverwijding optreedt. Een implicatie van deze bevinding zou kunnen zijn dat antimigraine middelen in de toekomst geen vaatvernauwend effect hoeven te hebben.

CONCLUSIES EN TOEKOMSTPERSPECTIEVEN

Gebaseerd op de bevindingen gepresenteerd in dit proefschrift kunnen er verschillende conclusies getrokken worden. De meest belangrijke is dat er geen vaatverwijding van hersenbloedvaten optreedt tijdens de hoofdpijnfase van een migraineaanval. Bij de ontwikkeling van nieuwe antimigraine middelen dient men zich te focussen op non vasculaire mechanismen. De tweede conclusie is dat er geen duidelijke relatie is tussen het optreden van mentale stress en het ontstaan van een migraine aanval. Het advies aan patiënten om stressvolle situaties te vermijden lijkt dan ook geen zinvol advies. De derde conclusie is dat hypoxie, zoals dit voorkomt in het hooggebergte, mogelijk een uitlokkende factor is voor migraine mogelijk via de ontwikkeling van cerebraal oedeem. Deze conclusie is echter behoorlijk speculatief en dient verder bestudeerd te worden.

REFERENCES

1. Menken M, Munsat TL, Toole JF. The global burden of disease study: implications for neurology. *Arch Neurol.* 2000; 57:418-420
2. Lipton RB, Stewart WF, Scher AI. Epidemiology and economic impact of migraine. *Curr Med Res Opin.* 2001; 17 Suppl 1:s4-12
3. Headache Classification Committee of the International Headache Society. The International Classification of Headache Disorders, 2nd edition. *Cephalalgia.* 2004; 24:1-160
4. Silberstein SD, Young WB. Migraine aura and prodrome. *Semin Neurol.* 1995; 15:175-182
5. Blau JN. Migraine prodromes separated from the aura: complete migraine. *Br Med J.* 1980; 281:658-660
6. Russell MB, Rasmussen BK, Fenger K et al. Migraine without aura and migraine with aura are distinct clinical entities: a study of four hundred and eighty-four male and female migraineurs from the general population. *Cephalalgia.* 1996; 16:239-245
7. Quintela E, Castillo J, Munoz P et al. Premonitory and resolution symptoms in migraine: a prospective study in 100 unselected patients. *Cephalalgia.* 2006; 26:1051-1060
8. Launer LJ, Terwindt GM, Ferrari MD. The prevalence and characteristics of migraine in a population-based cohort. The GEM study. *Neurology.* 1999; 53:537-542
9. Blumenthal HJ, Rapoport AM. The clinical spectrum of migraine. *Med Clin North Am.* 2001; 85:897-909
10. Lipton RB, Scher AI, Kolodner K et al. Migraine in the United States: epidemiology and patterns of health care use. *Neurology.* 2002; 58:885-894
11. Ferrari MD. Migraine. *Lancet.* 1998; 351:1043-1051
12. Pietrobon D, Striessnig J. Neurological diseases: Neurobiology of migraine. *Nat Rev Neurosci.* 2003; 4:386-398
13. Russell MB, Olesen J. Increased familial risk and evidence of genetic factor in migraine. *BMJ.* 1995; 311:541-544
14. Silberstein SD. Preventive treatment of headaches. *Curr Opin Neurol.* 2005; 18:289-292
15. Ophoff RA, Terwindt GM, Vergouwe MN et al. Familial Hemiplegic Migraine and Episodic Ataxia Type-2 Are Caused by Mutations in the CA2+ Channel Gene CACNL1A4. *Cell.* 1996; 87:543-552
16. De Fusco M, Marconi R, Silvestri L et al. Haploinsufficiency of ATP1A2 encoding the Na+/K+ pump alpha2 subunit associated with familial hemiplegic migraine type 2. *Nat Genet.* 2003; 33:192-196
17. Dichgans M, Freilinger T, Eckstein G et al. Mutation in the neuronal voltage-gated sodium channel SCN1A in familial hemiplegic migraine. *Lancet.* 2005; 366:371-377
18. Kors EE, Vanmolkot KR, Haan J et al. Recent findings in headache genetics. *Curr Opin Neurol.* 2004; 17:283-288

____ regel 1
____ regel 2
____ regel 3
____ regel 4
____ regel 5
____ regel 6
____ regel 7
____ regel 8
____ regel 9
____ regel 10
____ regel 11
____ regel 12
____ regel 13
____ regel 14
____ regel 15
____ regel 16
____ regel 17
____ regel 18
____ regel 19
____ regel 20
____ regel 21
____ regel 22
____ regel 23
____ regel 24
____ regel 25
____ regel 26
____ regel 27
____ regel 28
____ regel 29
____ regel 30
____ regel 31
____ regel 32
____ regel 33
____ regel 34
____ regel 35
____ regel 36
____ regel 37
____ regel 38
____ regel 39

References

- regel 1 _____
- regel 2 _____
- regel 3 _____
- regel 4 _____
- regel 5 _____
- regel 6 _____
- regel 7 _____
- regel 8 _____
- regel 9 _____
- regel 10 _____
- regel 11 _____
- regel 12 _____
- regel 13 _____
- regel 14 _____
- regel 15 _____
- regel 16 _____
- regel 17 _____
- regel 18 _____
- regel 19 _____
- regel 20 _____
- regel 21 _____
- regel 22 _____
- regel 23 _____
- regel 24 _____
- regel 25 _____
- regel 26 _____
- regel 27 _____
- regel 28 _____
- regel 29 _____
- regel 30 _____
- regel 31 _____
- regel 32 _____
- regel 33 _____
- regel 34 _____
- regel 35 _____
- regel 36 _____
- regel 37 _____
- regel 38 _____
- regel 39 _____
19. Kirchmann M, Thomsen LL, Olesen J. The CACNA1A and ATP1A2 genes are not involved in dominantly inherited migraine with aura. *Am J Med Genet B Neuropsychiatr Genet.* 2006; 141:250-256
 20. Martin VT, Behbehani MM. Toward a rational understanding of migraine trigger factors. *Med Clin North Am.* 2001; 85:911-941
 21. Lipton RB. Fair winds and foul headaches: risk factors and triggers of migraine. *Neurology.* 2000; 54:280-281
 22. Grant EC. Food allergies and migraine. *Lancet.* 1979; 1:966-969
 23. Van dB, V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache.* 1987; 27:191-196
 24. Takeshima T, Ishizaki K, Fukuhara Y et al. Population-based door-to-door survey of migraine in Japan: the Daisen study. *Headache.* 2004; 44:8-19
 25. Robbins L. Precipitating factors in migraine: a retrospective review of 494 patients. *Headache.* 1994; 34:214-216
 26. Rasmussen BK. Migraine and tension-type headache in a general population precipitating factors, femal hormones, sleep pattern and relation to lifestyle. *Pain.* 1993; 53:65-76
 27. Chabriat H, Dancho J, Michel P et al. Precipitating factors of headache. A prospective study in a national control-matched survey in migraineurs and nonmigraineurs. *Headache.* 1999; 39:335-338
 28. Turner LC, Molgaard CA, Gardner CH et al. Migraine trigger factors in non-clinical Mexican-American population in San Diego county: implications for etiology. *Cephalalgia.* 1995; 15:523-530
 29. Zivadinov R, Willheim K, Sepic-Grahovac D et al. Migraine and tension-type headache in Croatia: a population-based survey of precipitating factors. *Cephalalgia.* 2003; 23:336-343
 30. Koseoglu E, Nacar M, Talaslioglu A et al. Epidemiological and clinical characteristics of migraine and tension type headache in 1146 females in Kayseri, Turkey. *Cephalalgia.* 2003; 23:381-388
 31. Bic Z, Blix GG, Hopp HP et al. The influence of a low-fat diet on incidence and severity of migraine headaches. *J Womens Health Gend Based Med.* 1999; 8:623-630
 32. Egger J, Carter CM, Wilson J et al. Is migraine food allergy? A double-blind controlled trial of oligoantigenic diet treatment. *Lancet.* 1983; 2:865-869
 33. Medina JL, Diamond S. The role of diet in migraine. *Headache.* 1978; 18:31-34
 34. Littlewood JT, Gibb C, Glover V et al. Red wine as a cause of migraine. *Lancet.* 1988; 1:558-559
 35. Gibb CM, Davies PT, Glover V et al. Chocolate is a migraine-provoking agent. *Cephalalgia.* 1991; 11:93-95
 36. Marcus DA, Scharff L, Turk D et al. A double-blind provocative study of chocolate as a trigger of headache. *Cephalalgia.* 1997; 17:855-862
 37. Ziegler DK, Stewart R. Failure of tyramine to induce migraine. *Neurology.* 1977; 27:725-726
 38. Herbert J. Fortnightly review. Stress, the brain, and mental illness. *BMJ.* 1997; 315:530-535
 39. Mohr DC, Hart SL, Julian L et al. Association between stressful life events and exacerbation in multiple sclerosis: a meta-analysis. *BMJ.* 2004; 328:731

40. Chen E, Hanson MD, Paterson LQ et al. Socioeconomic status and inflammatory processes in childhood asthma: the role of psychological stress. *J Allergy Clin Immunol.* 2006; 117:1014-1020 _____ regel 1
41. Chandola T, Brunner E, Marmot M. Chronic stress at work and the metabolic syndrome: prospective study. *BMJ.* 2006; 332:521-525 _____ regel 2
42. Sorbi MJ, Maassen GH, Spierings EL. A time series analysis of daily hassles and mood changes in the 3 days before the migraine attack. *Behav Med.* 1996; 22:103-113 _____ regel 3
43. Holm JE, Lokken C, Myers TC. Migraine and stress: a daily examination of temporal relationships in women migraineurs. *Headache.* 1997; 37:553-558 _____ regel 4
44. Thomsen LL, Iversen HK, Boesen F et al. Transcranial Doppler and cardiovascular responses during cardiovascular autonomic tests in migraineurs during and outside attacks. *Brain.* 1995; 118:1319-1327 _____ regel 5
45. Havanka-Kanniainen H. Cardiovascular reflex responses during migraine attack. *Headache.* 1986; 26:442-446 _____ regel 6
46. Sances G, Granella F, Nappi R et al. Course of migraine during pregnancy and postpartum: a prospective study. *Cephalalgia.* 2003; 23:197-205 _____ regel 7
47. MacGregor EA. Oestrogen and attacks of migraine with and without aura. *Lancet Neurol.* 2004; 3:354-361 _____ regel 8
48. Kovats RS, Bouma MJ, Hajat S et al. El Nino and health. *Lancet.* 2003; 362:1481-1489 _____ regel 9
49. von Mackensen S, Hoeppe P, Maarouf A et al. Prevalence of weather sensitivity in Germany and Canada. *Int J Biometeorol.* 2005; 49:156-166 _____ regel 10
50. Larmande P, Hubert B, Sorabella A et al. [Influence of changes in climate and the calendar on the onset of a migraine crisis]. *Rev Neurol (Paris).* 1996; 152:38-43 _____ regel 11
51. De Matteis G, Vellante M, Marrelli A et al. Geomagnetic activity, humidity, temperature and headache: is there any correlation? *Headache.* 1994; 34:41-43 _____ regel 12
52. Villeneuve PJ, Szyszkowicz M, Stieb D et al. Weather and emergency room visits for migraine headaches in Ottawa, Canada. *Headache.* 2006; 46:64-72 _____ regel 13
53. Prince PB, Rapoport AM, Sheftell FD et al. The effect of weather on headache. *Headache.* 2004; 44:596-602 _____ regel 14
54. Cooke LJ, Rose MS, Becker WJ. Chinook winds and migraine headache. *Neurology.* 2000; 54:302-307 _____ regel 15
55. Sances G, Tassorelli C, Pucci E et al. Reliability of the nitroglycerin provocative test in the diagnosis of neurovascular headaches. *Cephalalgia.* 2004; 24:110-119 _____ regel 16
56. Afridi SK, Kaube H, Goadsby PJ. Glyceryl trinitrate triggers premonitory symptoms in migraineurs. *Pain.* 2004; 110:675-680 _____ regel 17
57. Briganti A, Salonia A, Deho F et al. Clinical update on phosphodiesterase type-5 inhibitors for erectile dysfunction. *World J Urol.* 2005; 23:374-384 _____ regel 18
58. Kruuse C, Thomsen LL, Birk S et al. Migraine can be induced by sildenafil without changes in middle cerebral artery diameter. *Brain.* 2003; 126:241-247 _____ regel 19

References

- regel 1 _____ 59. Leone M, Attanasio A, Croci D et al. The serotonergic agent m-chlorophenylpiperazine induces migraine attacks: A controlled study. *Neurology*. 2000; 55:136-139
- regel 2 _____ 60. Brewerton TD, Murphy DL, Mueller EA et al. Introduction of migraine like headache by the serotonin agonist m-chlorophenylpiperazine. *Clin Pharmacol Ther*. 1988; 43:605-609
- regel 3 _____ 61. Goadsby PJ, Edvinsson L, Ekman R. Vasoactive peptide release in the extracerebral circulation of humans during migraine headache. *Ann Neurol*. 1990; 28:183-187
- regel 4 _____ 62. Lassen LH, Haderslev PA, Jacobson VB et al. CGRP may play a causative role in migraine. *Cephalalgia*. 2002; 22:54-61
- regel 5 _____ 63. Krabbe AA, Olesen J. Headache provocation by continuous intravenous infusion of histamine. Clinical results and receptor mechanisms. *Pain*. 1980; 8:253-259
- regel 6 _____ 64. Lassen LH, Thomsen LL, Olesen J. Histamine induces migraine via the H1-receptor. Support for the NO hypothesis of migraine. *NeuroReport*. 1995; 6:1475-1479
- regel 7 _____ 65. Shirai T, Meyer JS, Akiyama H et al. Acetazolamide testing of cerebral vasodilator capacity provokes "vascular" but not tension headaches. *Headache*. 1996; 36:589-594
- regel 8 _____ 66. Haan J, Sluis P, Sluis LH et al. Acetazolamide treatment for migraine aura status [In Process Citation]. *Neurology*. 2000; 55:1588-1589
- regel 9 _____ 67. Vahedi K, Taupin P, Djomby R et al. Efficacy and tolerability of acetazolamide in migraine prophylaxis: a randomised placebo-controlled trial. *J Neurol*. 2002; 249:206-211
- regel 10 _____ 68. Carlson LA, Ekelund LG, Oro L. Clinical and metabolic effects of different doses of prostaglandin E1 in man. Prostaglandin and related factors. *Acta Med Scand*. 1968; 183:423-430
- regel 11 _____ 69. Carroll JD, Hilton BP. The effects of reserpine injection on methysergide treated control and migrainous subjects. *Headache*. 1974; 14:149-156
- regel 12 _____ 70. Ferrari U, Empl M, Kim KS et al. Calcineurin inhibitor-induced headache: clinical characteristics and possible mechanisms. *Headache*. 2005; 45:211-214
- regel 13 _____ 71. Ratinahirana H, Benigni JP, Bousser MG. Injection of polidocanol foam (PF) in varicose veins as a trigger for attacks of migraine with visual aura. *Cephalalgia*. 2003; 23:850-851
- regel 14 _____ 72. Ramadan NM, Gilkey SJ, Mitchell M et al. Postangiography headache. *Headache*. 1995; 35:21-24
- regel 15 _____ 73. Cao Y, Aurora SK, Nagesh V et al. Functional MRI-BOLD of brainstem structures during visually triggered migraine. *Neurology*. 2002; 59:72-78
- regel 16 _____ 74. Frese A, Eikermann A, Frese K et al. Headache associated with sexual activity: demography, clinical features, and comorbidity. *Neurology*. 2003; 61:796-800
- regel 17 _____ 75. Goadsby PJ, Lipton RB, Ferrari MD. Migraine--current understanding and treatment. *N Engl J Med*. 2002; 346:257-270
- regel 18 _____ 76. Edvinsson L. Correlation between CGRP and migraine attacks. *Cephalalgia*. 2005; 25:163-164
- regel 19 _____ 77. Bolay H, Reuter U, Dunn AK et al. Intrinsic brain activity triggers trigeminal meningeal afferents in a migraine model. *Nat Med*. 2002; 8:136-142
- regel 20 _____
- regel 21 _____
- regel 22 _____
- regel 23 _____
- regel 24 _____
- regel 25 _____
- regel 26 _____
- regel 27 _____
- regel 28 _____
- regel 29 _____
- regel 30 _____
- regel 31 _____
- regel 32 _____
- regel 33 _____
- regel 34 _____
- regel 35 _____
- regel 36 _____
- regel 37 _____
- regel 38 _____
- regel 39 _____

78. Thomsen LL, Kruuse C, Iversen HK et al. A nitric oxide donor (nitroglycerin) triggers genuine migraine attacks. *Eur J Neurol.* 1994; 1:73-80 _____ regel 1
79. Waeber C, Moskowitz MA. Migraine as an inflammatory disorder. *Neurology.* 2005; 64:59-15 _____ regel 2
80. Pascual J, Arco del C, Romon T et al. Autoradiographic distribution of [3H]sumatriptan-binding sites in post-mortem human brain. *Cephalalgia.* 1996; 16:317-322 _____ regel 3
81. Knight YE. Brainstem modulation of caudal trigeminal nucleus: a model for understanding migraine biology and future drug targets. *Headache Currents.* 2005; 2:108-118 _____ regel 4
82. Weiller C, May A, Limmroth V et al. Brain stem activation in spontaneous human migraine attacks. *Nat Med.* 1995; 1:658-661 _____ regel 5
83. Afridi SK, Matharu MS, Lee L et al. A PET study exploring the laterality of brainstem activation in migraine using glyceryl trinitrate. *Brain.* 2005; _____ regel 6
84. Montagna P. Hypothalamus, sleep and headaches. *Neurol Sci.* 2006; 27 Suppl 2:S138-S143 _____ regel 7
85. Overeem S, van Vliet JA, Lammers GJ et al. The hypothalamus in episodic brain disorders. *Lancet Neurol.* 2002; 1:437-444 _____ regel 8
86. May A, Bahra A, Buchel C et al. Hypothalamic activation in cluster headache attacks. *Lancet.* 1998; 352:275-278 _____ regel 9
87. Parsons AA, Strijbos PJ. The neuronal versus vascular hypothesis of migraine and cortical spreading depression. *Curr Opin Pharmacol.* 2003; 3:73-77 _____ regel 10
88. Fanciullacci C, Alessandri M, Fanciullacci M. The relationship between stress and migraine. *Funct Neurol.* 1998; 13:215-223 _____ regel 11
89. Van den Bergh V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache.* 1987; 27:191-196 _____ regel 12
90. Wittrock DA, Foraker SL. Tension-type headache and stressful events: the role of selective memory in the reporting of stressors. *Headache.* 2001; 41:482-493 _____ regel 13
91. Ziegler DK, Hassanein RS, Kodanaz A et al. Circadian rhythms of plasma cortisol in migraine. *J Neurol Neurosurg Psychiatry.* 1979; 42:741-748 _____ regel 14
92. van Hilten JJ, Ferrari MD, Van der Meer JW et al. Plasma interleukin-1, tumour necrosis factor and hypothalamic-pituitary- adrenal axis responses during migraine attacks. *Cephalalgia.* 1991; 11:65-67 _____ regel 15
93. Avnon Y, Nitzan M, Sprecher E et al. Autonomic asymmetry in migraine: augmented parasympathetic activation in left unilateral migraineurs. *Brain.* 2004; 127:2099-2108 _____ regel 16
94. Takeshima T, Takao Y, Takahashi K. Pupillary sympathetic hypofunction and asymmetry in muscle contraction headache and migraine. *Cephalalgia.* 1987; 7:257-262 _____ regel 17
95. Shechter A, Stewart WF, Silberstein SD et al. Migraine and autonomic nervous system function: A population-based, case-control study. *Neurology.* 2002; 58:422-427 _____ regel 18
96. Hassinger HJ, Semenchuk EM, O'Brien WH. Cardiovascular responses to pain and stress in migraine. *Headache.* 1999; 39:605-615 _____ regel 19
97. Hackett PH, Roach RC. High-altitude illness. *N Engl J Med.* 2001; 345:107-114 _____ regel 20

References

- regel 1 _____ 98. Appenzeller O. Altitude headache. *Headache*. 1972; 12:126-129
- regel 2 _____ 99. Schneider, M, Bernasch, D, Weymann, J, and Bartsch, P. Characteristics of high altitude headache. *High Alt Med Biol Abstract Vth World Congress on Mountain Medicin Barcelona*[3], 100. 2002. Ref Type: Abstract
- regel 3 _____
- regel 4 _____
- regel 5 _____ 100. Arregui A, Leon-Velarde F, Cabrera J et al. Migraine, polycythemia and chronic mountain sickness. *Cephalalgia*. 1994; 14:339-341
- regel 6 _____
- regel 7 _____ 101. Jaillard AS, Mazetti P, Kala E. Prevalence of migraine and headache in a high-altitude town of Peru: a population-based study. *Headache*. 1997; 37:95-101
- regel 8 _____
- regel 9 _____ 102. Bartsch P, Maggi S, Kleger GR et al. Sumatriptan for high-altitude headache. *Lancet*. 1994; 344:1445
- regel 10 _____ 103. Burtscher M, Likar R, Nachbauer W et al. Ibuprofen versus sumatriptan for high-altitude headache. *Lancet*. 1995; 346:254-255
- regel 11 _____
- regel 12 _____ 104. Hackett PH, Yarnell PR, Hill R et al. High-altitude cerebral edema evaluated with magnetic resonance imaging: clinical correlation and pathophysiology. *JAMA*. 1998; 280:1920-1925
- regel 13 _____
- regel 14 _____ 105. Bates D, Ashford E, Dawson R et al. Subcutaneous sumatriptan during migraine aura. *Neurology*. 1994; 44:1587-1592
- regel 15 _____
- regel 16 _____ 106. Moncada S, Palmer RM, Higgs EA. Nitric oxide: physiology, pathophysiology, and pharmacology. *Pharmacol Rev*. 1991; 43:109-142
- regel 17 _____
- regel 18 _____ 107. Tassorelli C, Joseph SA, Buzzi MG et al. The effects on the central nervous system of nitroglycerin-- putative mechanisms and mediators. *Prog Neurobiol*. 1999; 57:607-624
- regel 19 _____
- regel 20 _____ 108. Olesen J, Thomsen LL, Lassen LH et al. The nitric oxide hypothesis of migraine and other vascular headaches. *Cephalalgia*. 1995; 15(2):94-100
- regel 21 _____
- regel 22 _____ 109. Hansen J, Pedersen D, Larsen V et al. Magnetic resonance angiography shows dilatation of the middle cerebral artery after infusion of glyceryl trinitrate in healthy volunteers. *Cephalalgia*. 2007; 27:118-127
- regel 23 _____
- regel 24 _____ 110. Schoonman GG, Bakker D, Schmitz N et al. Magnetic resonance angiography of the human middle meningeal artery: Implications for migraine. *J Magn Reson Imaging*. 2006; 24:918-921
- regel 25 _____
- regel 26 _____ 111. Reinert M, Wiest R, Barth L et al. Transdermal nitroglycerin in patients with subarachnoid hemorrhage. *Neurol Res*. 2004; 26:435-439
- regel 27 _____
- regel 28 _____ 112. Dahl A, Russell D, Nyberg-Hansen R et al. Effect of nitroglycerin on cerebral circulation measured by transcranial Doppler and SPECT. *Stroke*. 1989; 20:1733-1736
- regel 29 _____
- regel 30 _____ 113. White RP, Deane C, Hindley C et al. The effect of the nitric oxide donor glyceryl trinitrate on global and regional cerebral blood flow in man. *J Neurol Sci*. 2000; 178:23-28
- regel 31 _____
- regel 32 _____ 114. Wei EP, Moskowitz MA, Boccalini P et al. Calcitonin gene-related peptide mediates nitroglycerin and sodium nitroprusside-induced vasodilation in feline cerebral arterioles. *Circ Res*. 1992; 70:1313-1319
- regel 33 _____
- regel 34 _____ 115. Zhou ZH, Deng HW, Li YJ. The depressor effect of nitroglycerin is mediated by calcitonin gene-related peptide. *Life Sci*. 2001; 69:1313-1320
- regel 35 _____
- regel 36 _____ 116. Sicuteri F, Del Bene E, Poggioni M et al. Unmasking latent dysnociception in healthy subjects. *Headache*. 1987; 27:180-185
- regel 37 _____
- regel 38 _____
- regel 39 _____

117. Thomsen LL, Iversen HK, Brinck TA et al. Arterial supersensitivity to nitric oxide (nitroglycerin) in migraine sufferers. *Cephalalgia*. 1993; 13:395-399 — regel 1
118. Zanette EM, Agnoli A, Cerbo R et al. Transcranial doppler (TCD) after nitroglycerin in migraine without aura. *Headache*. 1991; 31:596-598 — regel 2
119. Van dB, V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache*. 1987; 27:191-196 — regel 3
120. Van dB, V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache*. 1987; 27:191-196 — regel 4
121. Van dB, V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache*. 1987; 27:191-196 — regel 5
122. Van dB, V, Amery WK, Waelkens J. Trigger factors in migraine: a study conducted by the Belgian Migraine Society. *Headache*. 1987; 27:191-196 — regel 6
123. Tvedskov J, Iversen H, Olesen J. A double-blind study of SB-220453 (Tonerbasat) in the glyceryltrinitrate (GTN) model of migraine. *Cephalalgia*. 2004; 24:875-882 — regel 7
124. Juhasz G, Zsombok T, Modos EA et al. NO-induced migraine attack: strong increase in plasma calcitonin gene-related peptide (CGRP) concentration and negative correlation with platelet serotonin release. *Pain*. 2003; 106:461-470 — regel 8
125. Bellantonio P, Micieli G, Buzzi MG et al. Haemodynamic correlates of early and delayed responses to sublingual administration of isosorbide dinitrate in migraine patients: a transcranial Doppler study. *Cephalalgia*. 1997; 17:183-187 — regel 9
126. Schoonman G, Sandor P, Agosti R et al. Normobaric hypoxia and nitroglycerin as trigger factors for migraine. *Cephalalgia*. 2006; 26:816-819 — regel 10
127. Tvedskov JF, Thomsen LL, Iversen HK et al. The prophylactic effect of valproate on glyceryltrinitrate induced migraine. *Cephalalgia*. 2004; 24:576-585 — regel 11
128. Juhasz G, Zsombok T, Jakab B et al. Sumatriptan causes parallel decrease in plasma calcitonin gene-related peptide (CGRP) concentration and migraine headache during nitroglycerin induced migraine attack. *Cephalalgia*. 2005; 25:179-183 — regel 12
129. Tvedskov JF, Thomsen LL, Thomsen LL et al. The effect of propranolol on glyceryltrinitrate-induced headache and arterial response. *Cephalalgia*. 2004; 24:1076-1087 — regel 13
130. Kruuse C, Lassen LH, Iversen HK et al. Dipyridamole may induce migraine in patients with migraine without aura. *Cephalalgia*. 2006; 26:925-933 — regel 14
131. Giffin NJ, Ruggiero L, Lipton RB et al. Premonitory symptoms in migraine: An electronic diary study. *Neurology*. 2003; 60:935-940 — regel 15
132. Rasmussen BK, Olesen J. Migraine with aura and migraine without aura: an epidemiological study. *Cephalalgia*. 1992; 12:221-228 — regel 16
133. Amery WK, Waelkens J, Vandenbergh V. Migraine warnings. *Headache*. 1986; 26:60-66 — regel 17

References

- regel 1 _____ 134. Kelman L. The premonitory symptoms (prodrome): a tertiary care study of 893 migraineurs. *Headache*. 2004; 44:865-872
- regel 2 _____
- regel 3 _____ 135. Santoro G, Bernasconi F, Sessa F et al. Premonitory symptoms in migraine without aura: a clinical investigation. *Funct Neurol*. 1990; 5:339-344
- regel 4 _____
- regel 5 _____ 136. Freeman EW. Premenstrual syndrome and premenstrual dysphoric disorder: definitions and diagnosis. *Psychoneuroendocrinology*. 2003;25-37
- regel 6 _____
- regel 7 _____ 137. Zellner DA, Garriga-Trillo A, Rohm E et al. Food liking and craving: A cross-cultural approach. *Appetite*. 1999; 33:61-70
- regel 8 _____
- regel 9 _____ 138. Chen W, Woods SL, Puntillo KA. Gender differences in symptoms associated with acute myocardial infarction: a review of the research. *Heart Lung*. 2005; 34:240-247
- regel 10 _____
- regel 11 _____ 139. Kogan A, Eidelman LA, Raanani E et al. Nausea and vomiting after fast-track cardiac anaesthesia. *Br J Anaesth*. 2003; 91:214-217
- regel 12 _____
- regel 13 _____ 140. American Psychiatric Association. *Diagnostic and statistical manual of mental disorders*. 4th rev. ed. Washington, DC: American Psychiatric Press, 1994
- regel 14 _____
- regel 15 _____ 141. Passchier J. A critical note on psychophysiological stress research into migraine patients. *Cephalalgia*. 1994; 14:194-198
- regel 16 _____
- regel 17 _____ 142. Loong TW. Understanding sensitivity and specificity with the right side of the brain. *BMJ*. 2003; 327:716-719
- regel 18 _____
- regel 19 _____ 143. de Kloet ER, Joels M, Holsboer F. Stress and the brain: from adaptation to disease. *Nat Rev Neurosci*. 2005; 6:463-475
- regel 20 _____
- regel 21 _____ 144. Cohen S, Kamarck T, Mermelstein R. A global measure of perceived stress. *J Health Soc Behav*. 1983; 24:385-396
- regel 22 _____
- regel 23 _____ 145. Meyer TJ, Miller ML, Metzger RL et al. Development and validation of the Penn State Worry Questionnaire. *Behav Res Ther*. 1990; 28:487-495
- regel 24 _____
- regel 25 _____ 146. Laskin RS, Davis JP. The use of a personal digital assistant in orthopaedic surgical practice. *Clin Orthop*. 2004;91-98
- regel 26 _____
- regel 27 _____ 147. Brantley PJ, Waggoner CD, Jones GN et al. A Daily Stress Inventory: development, reliability, and validity. *J Behav Med*. 1987; 10:61-74
- regel 28 _____
- regel 29 _____ 148. van Aken MO, Romijn JA, Miltenburg JA et al. Automated measurement of salivary cortisol. *Clin Chem*. 2003; 49:1408-1409
- regel 30 _____
- regel 31 _____ 149. de Geus EJ, Willemsen GH, Klaver CH et al. Ambulatory measurement of respiratory sinus arrhythmia and respiration rate. *Biol Psychol*. 1995; 41:205-227
- regel 32 _____
- regel 33 _____ 150. Bilchick KC, Berger RD. Heart rate variability. *J Cardiovasc Electrophysiol*. 2006; 17:691-694
- regel 34 _____
- regel 35 _____ 151. Havanka-Kannianen H, Tolonen U, Myllyla VV. Cardiovascular reflexes in young migraine patients. *Headache*. 1986; 26:420-424
- regel 36 _____
- regel 37 _____ 152. Avnon Y, Nitzan M, Sprecher E et al. Different patterns of parasympathetic activation in uni- and bilateral migraineurs. *Brain*. 2003; 126:1660-1670
- regel 38 _____
- regel 39 _____

153. Smyth J, Ockenfels MC, Porter L et al. Stressors and mood measured on a momentary basis are associated with salivary cortisol secretion. *Psychoneuroendocrinology*. 1998; 23:353-370 _____ regel 1
154. Brantley PJ, Dietz LS, McKnight GT et al. Convergence between the Daily Stress Inventory and endocrine measures of stress. *J Consult Clin Psychol*. 1988; 56:549-551 _____ regel 2
155. Vrijkotte TG, van Doornen LJ, de Geus EJ. Effects of work stress on ambulatory blood pressure, heart rate, and heart rate variability. *Hypertension*. 2000; 35:880-886 _____ regel 3
156. Bruni O, Russo PM, Violani C et al. Sleep and migraine: an actigraphic study. *Cephalalgia*. 2004; 24:134-139 _____ regel 4
157. Kirschbaum C, Kudielka BM, Gaab J et al. Impact of gender, menstrual cycle phase, and oral contraceptives on the activity of the hypothalamus-pituitary-adrenal axis. *Psychosom Med*. 1999; 61:154-162 _____ regel 5
158. Lundberg U, Hellstrom b. Workload and morning salivary cortisol in women. *Work & Stress*. 2002; 16:356-363 _____ regel 6
159. Deinzer R, Kirschbaum C, Gresele C et al. Adrenocortical responses to repeated parachute jumping and subsequent h- CRH challenge in inexperienced healthy subjects. *Physiol Behav*. 1997; 61:507-511 _____ regel 7
160. Lundberg U. Stresshormones in health and illness: the roles of work and gender. *Psychoneuroendocrinology*. 2005; 30:1017-1021 _____ regel 8
161. James GD, Berge-Landry HH, Valdimarsdottir HB et al. Urinary catecholamine levels in daily life are elevated in women at familial risk of breast cancer. *Psychoneuroendocrinology*. 2004; 29:831-838 _____ regel 9
162. Christiansen I, Thomsen LL, Daugaard D et al. Glyceryl trinitrate induces attacks of migraine without aura in sufferers of migraine with aura. *Cephalalgia*. 1999; 19:660-667 _____ regel 10
163. Roach RC, Bartsch P, Hackett PH et al. The Lake Louise AMS Scoring Consensus Committee. The Lake Louise acute mountain sickness scoring system. Burlington: Queen City Printers Inc, 1993 _____ regel 11
164. Basnyat B, Murdoch DR. High-altitude illness. *Lancet*. 2003; 361:1967-1974 _____ regel 12
165. Levine BD, Yoshimura K, Kobayashi T et al. Dexamethasone in the treatment of acute mountain sickness. *N Engl J Med*. 1989; 321:1707-1713 _____ regel 13
166. Sutton JR, Lassen N. Pathophysiology of acute mountain sickness and high altitude pulmonary oedema: an hypothesis. *Bull Physiopathol Respir (Nancy)*. 1979; 15:1045-1052 _____ regel 14
167. Kallenberg K, Bailey DM, Christ S et al. Magnetic resonance imaging evidence of cytotoxic cerebral edema in acute mountain sickness. *J Cereb Blood Flow Metab*. 2006; _____ regel 15
168. Morocz IA, Zientara GP, Gudbjartsson H et al. Volumetric quantification of brain swelling after hypobaric hypoxia exposure. *Exp Neurol*. 2001; 168:96-104 _____ regel 16
169. Fischer R, Vollmar C, Thiere M et al. No evidence of cerebral oedema in severe acute mountain sickness. *Cephalalgia*. 2004; 24:66-71 _____ regel 17
170. Matsuzawa YK, Kobayashi T, Fujimoto K et al. Cerebral edema in acute mountain sickness. In: Ueda G, Reeves JT, Sekiguchi M, eds. *High Altitude Medicine*., 1992:300-304 _____ regel 18

References

- regel 1 _____ 171. Sampson JB, Cymerman A, Burse RL et al. Procedures for the measurement of acute mountain sickness. *Aviat Space Environ Med.* 1983; 54:1063-1073
- regel 2 _____
- regel 3 _____ 172. Patel MR, Siewert B, Warach S et al. Diffusion and perfusion imaging techniques. *Magn Reson Imaging Clin N Am.* 1995; 3:425-438
- regel 4 _____
- regel 5 _____ 173. Mintorovitch J, Moseley ME, Chileuitt L et al. Comparison of diffusion- and T2-weighted MRI for the early detection of cerebral ischemia and reperfusion in rats. *Magn Reson Med.* 1991; 18:39-50
- regel 6 _____
- regel 7 _____ 174. Kucharczyk J, Vexler ZS, Roberts TP et al. Echo-planar perfusion-sensitive MR imaging of acute cerebral ischemia. *Radiology.* 1993; 188:711-717
- regel 8 _____
- regel 9 _____ 175. Mintorovitch J, Yang GY, Shimizu H et al. Diffusion-weighted magnetic resonance imaging of acute focal cerebral ischemia: comparison of signal intensity with changes in brain water and Na⁺,K⁽⁺⁾-ATPase activity. *J Cereb Blood Flow Metab.* 1994; 14:332-336
- regel 10 _____
- regel 11 _____
- regel 12 _____ 176. Brunberg JA, Chenevert TL, McKeever PE et al. In vivo MR determination of water diffusion coefficients and diffusion anisotropy: correlation with structural alteration in gliomas of the cerebral hemispheres. *AJNR Am J Neuroradiol.* 1995; 16:361-371
- regel 13 _____
- regel 14 _____
- regel 15 _____ 177. Tien RD, Felsberg GJ, Friedman H et al. MR imaging of high-grade cerebral gliomas: value of diffusion-weighted echoplanar pulse sequences. *AJR Am J Roentgenol.* 1994; 162:671-677
- regel 16 _____
- regel 17 _____ 178. Bartsch P, Baumgartner RW, Waber U et al. Comparison of carbon-dioxide-enriched, oxygen-enriched, and normal air in treatment of acute mountain sickness. *Lancet.* 1990; 336:772-775
- regel 18 _____
- regel 19 _____ 179. Roach RC, Loeppky JA, Icenogle MV. Acute mountain sickness: increased severity during simulated altitude compared with normobaric hypoxia. *J Appl Physiol.* 1996; 81:1908-1910
- regel 20 _____
- regel 21 _____ 180. Scarabino T, Nemore F, Giannatempo GM et al. 3.0 T magnetic resonance in neuroradiology. *Eur J Radiol.* 2003; 48:154-164
- regel 22 _____
- regel 23 _____ 181. Ogawa S, Lee TM, Kay AR et al. Brain magnetic resonance imaging with contrast dependent on blood oxygenation. *Proc Natl Acad Sci U S A.* 1990; 87:9868-9872
- regel 24 _____
- regel 25 _____ 182. Ogawa S, Lee TM, Nayak AS et al. Oxygenation-sensitive contrast in magnetic resonance image of rodent brain at high magnetic fields. *Magn Reson Med.* 1990; 14:68-78
- regel 26 _____
- regel 27 _____ 183. Prielmeier F, Nagatomo Y, Frahm J. Cerebral blood oxygenation in rat brain during hypoxic hypoxia. Quantitative MRI of effective transverse relaxation rates. *Magn Reson Med.* 1994; 31:678-681
- regel 28 _____
- regel 29 _____ 184. Lin W, Paczynski RP, Celik A et al. Experimental hypoxemic hypoxia: changes in R2* of brain parenchyma accurately reflect the combined effects of changes in arterial and cerebral venous oxygen saturation. *Magn Reson Med.* 1998; 39:474-481
- regel 30 _____
- regel 31 _____
- regel 32 _____ 185. van Zijl PC, Eleff SM, Ulatowski JA et al. Quantitative assessment of blood flow, blood volume and blood oxygenation effects in functional magnetic resonance imaging. *Nat Med.* 1998; 4:159-167
- regel 33 _____
- regel 34 _____ 186. Rostrup E, Larsson HB, Born AP et al. Changes in BOLD and ADC weighted imaging in acute hypoxia during sea-level and altitude adapted states. *Neuroimage.* 2005;
- regel 35 _____
- regel 36 _____ 187. Sanchez dR, Moskowitz MA. High altitude headache. Lessons from headaches at sea level. *Adv Exp Med Biol.* 1999; 474:145-153
- regel 37 _____
- regel 38 _____
- regel 39 _____

188. Ferrazzini G, Maggiorini M, Kriemler S et al. Successful treatment of acute mountain sickness with dexamethasone. *Br Med J (Clin Res Ed)*. 1987; 294:1380-1382 _____ regel 1
189. Allen K, Busza AL, Crockard HA et al. Brain metabolism and blood flow in acute cerebral hypoxia studied by NMR spectroscopy and hydrogen clearance. *NMR Biomed*. 1992; 5:48-52 _____ regel 2
190. De Angelis C, Hauptert GT, Jr. Hypoxia triggers release of an endogenous inhibitor of Na(+)-K(+)-ATPase from midbrain and adrenal. *Am J Physiol*. 1998; 274:F182-F188 _____ regel 3
191. Nioka S, Smith DS, Chance B et al. Oxidative phosphorylation system during steady-state hypoxia in the dog brain. *J Appl Physiol*. 1990; 68:2527-2535 _____ regel 4
192. Ben Yoseph O, Badar-Goffer RS, Morris PG et al. Glycerol 3-phosphate and lactate as indicators of the cerebral cytoplasmic redox state in severe and mild hypoxia respectively: a ¹³C- and ³¹P-n.m.r. study. *Biochem J*. 1993; 291 (Pt 3):915-919 _____ regel 5
193. Rolett EL, Azzawi A, Liu KJ et al. Critical oxygen tension in rat brain: a combined (³¹P)-NMR and EPR oximetry study. *Am J Physiol Regul Integr Comp Physiol*. 2000; 279:R9-R16 _____ regel 6
194. Baumgartner RW, Eichenberger U, Bartsch P. Postural ataxia at high altitude is not related to mild to moderate acute mountain sickness. *Eur J Appl Physiol*. 2002; 86:322-326 _____ regel 7
195. Moller K, Paulson OB, Hornbein TF et al. Unchanged cerebral blood flow and oxidative metabolism after acclimatization to high altitude. *J Cereb Blood Flow Metab*. 2002; 22:118-126 _____ regel 8
196. Severinghaus JW, Chiodi H, Eger EI et al. Cerebral blood flow in man at high altitude. Role of cerebrospinal fluid pH in normalization of flow in chronic hypocapnia. *Circ Res*. 1966; 19:274-282 _____ regel 9
197. Guadagno JV, Jones PS, Fryer TD et al. Local relationships between restricted water diffusion and oxygen consumption in the ischemic human brain. *Stroke*. 2006; 37:1741-1748 _____ regel 10
198. Wolff HG. Pain-sensitive structures within the cranial cavity. *Headache and other head pain.*, 1948:59-97 _____ regel 11
199. Ferrari MD, Roon KI, Lipton RB et al. Oral triptans (serotonin 5-HT(1B/1D) agonists) in acute migraine treatment: a meta-analysis of 53 trials. *Lancet*. 2001; 358:1668-1675 _____ regel 12
200. Henkes H, May A, Kuhne D et al. Sumatriptan: vasoactive effect on human dural vessels, demonstrated by subselective angiography. *Cephalalgia*. 1996; 16:224-230 _____ regel 13
201. Berlis A, Putz R, Schumacher M. Direct and CT measurements of canals and foramina of the skull base. *Br J Radiol*. 1992; 65:653-661 _____ regel 14
202. Bahra A, Matharu MS, Buchel C et al. Brainstem activation specific to migraine headache. *Lancet*. 2001; 357:1016-1017 _____ regel 15
203. van der Geest RJ, Reiber JH. Quantification in cardiac MRI. *J Magn Reson Imaging*. 1999; 10:602-608 _____ regel 16
204. Friberg L, Olesen J, Iversen HK et al. Migraine pain associated with middle cerebral artery dilatation: reversal by sumatriptan. *Lancet*. 1991; 338:13-17 _____ regel 17
205. Maier W, Windecker S, Kung A et al. Exercise-induced coronary artery vasodilation is not impaired by stent placement. *Circulation*. 2002; 105:2373-2377 _____ regel 18

References

- regel 1 _____ 206. Hoogeveen RM, Bakker CJ, Viergever MA. Limits to the accuracy of vessel diameter measurement in MR angiography. *J Magn Reson Imaging*. 1998; 8:1228-1235
- regel 2 _____
- regel 3 _____ 207. Bednarczyk EM, Wack DS, Kassab MY et al. Brain blood flow in the nitroglycerin (GTN) model of migraine: measurement using positron emission tomography and transcranial Doppler. *Cephalalgia*. 2002; 22:749-757
- regel 4 _____
- regel 5 _____
- regel 6 _____ 208. May A, Goadsby PJ. The trigeminovascular system in humans: pathophysiologic implications for primary headache syndromes of the neural influence on the cerebral circulation. *J Cereb Blood Flow Metab*. 1999; 19:115-127
- regel 7 _____
- regel 8 _____
- regel 9 _____ 209. Pluta RM. Delayed cerebral vasospasm and nitric oxide: review, new hypothesis, and proposed treatment. *Pharmacol Ther*. 2005; 105:23-56
- regel 10 _____
- regel 11 _____ 210. Thomsen LL. Investigations into the role of nitric oxide and the large intracranial arteries in migraine headache. *Cephalalgia*. 1997; 17:873-895
- regel 12 _____
- regel 13 _____ 211. Spilt A, Van den BR, Kamper AM et al. MR assessment of cerebral vascular response: a comparison of two methods. *J Magn Reson Imaging*. 2002; 16:610-616
- regel 14 _____
- regel 15 _____ 212. Spilt A, Box FM, van der Geest RJ et al. Reproducibility of total cerebral blood flow measurements using phase contrast magnetic resonance imaging. *J Magn Reson Imaging*. 2002; 16:1-5
- regel 16 _____
- regel 17 _____ 213. Bakker CJ, Hartkamp MJ, Mali WP. Measuring blood flow by nontriggered 2D phase-contrast MR angiography. *Magn Reson Imaging*. 1996; 14:609-614
- regel 18 _____
- regel 19 _____ 214. Bakker CJ, Kouwenhoven M, Hartkamp MJ et al. Accuracy and precision of time-averaged flow as measured by nontriggered 2D phase-contrast MR angiography, a phantom evaluation. *Magn Reson Imaging*. 1995; 13:959-965
- regel 20 _____
- regel 21 _____
- regel 22 _____ 215. de Koning PJ, Schaap JA, Janssen JP et al. Automated segmentation and analysis of vascular structures in magnetic resonance angiographic images. *Magn Reson Med*. 2003; 50:1189-1198
- regel 23 _____
- regel 24 _____ 216. Gisolf J, Westerhof BE, van Dijk N et al. Sublingual nitroglycerin used in routine tilt testing provokes a cardiac output-mediated vasovagal response. *J Am Coll Cardiol*. 2004; 44:588-593
- regel 25 _____
- regel 26 _____ 217. Hamel E. Perivascular nerves and the regulation of cerebrovascular tone. *J Appl Physiol*. 2006; 100:1059-1064
- regel 27 _____
- regel 28 _____ 218. Spilt A, Van den BR, Kamper AM et al. MR assessment of cerebral vascular response: a comparison of two methods. *J Magn Reson Imaging*. 2002; 16:610-616
- regel 29 _____
- regel 30 _____ 219. de Boorder MJ, Hendrikse J, van der GJ. Phase-contrast magnetic resonance imaging measurements of cerebral autoregulation with a breath-hold challenge: a feasibility study. *Stroke*. 2004; 35:1350-1354
- regel 31 _____
- regel 32 _____ 220. Vanninen E, Kuikka JT, Tenhunen-Eskelinen M et al. Haemodynamic effects of acetazolamide in patients with cardiovascular disorders: correlation with calculated cerebral perfusion reserve. *Nucl Med Commun*. 1996; 17:325-330
- regel 33 _____
- regel 34 _____
- regel 35 _____ 221. Ferrari MD, Saxena PR. Clinical and experimental effects of sumatriptan in humans. *Trends Pharmacol Sci*. 1993; 14:129-133
- regel 36 _____
- regel 37 _____
- regel 38 _____
- regel 39 _____

222. Nichols FT, Mawad M, Mohr JP et al. Focal headache during balloon inflation in the internal carotid and middle cerebral arteries. *Stroke*. 1990; 21:555-559 _____ regel 1
223. Olesen J, Diener HC, Husstedt IW et al. Calcitonin gene-related peptide receptor antagonist BIBN 4096 BS for the acute treatment of migraine. *N Engl J Med*. 2004; 350:1104-1110 _____ regel 2
224. Doods H, Arndt K, Rudolf K et al. CGRP antagonists: unravelling the role of CGRP in migraine. *Trends Pharmacol Sci*. 2007; 28:580-587 _____ regel 3
225. Tfelt-Hansen P, De Vries P, Saxena PR. Triptans in migraine: a comparative review of pharmacology, pharmacokinetics and efficacy. *Drugs*. 2000; 60:1259-1287 _____ regel 4
226. Iversen HK, Nielsen TH, Olesen J et al. Arterial responses during migraine headache. *Lancet*. 1990; 336:837-839 _____ regel 5
227. Thomsen LL, Iversen HK, Olesen J. Cerebral blood flow velocities are reduced during attacks of unilateral migraine without aura. *Cephalalgia*. 1995; 15:109-116 _____ regel 6
228. Edvinsson L, Uddman E, Wackenfors A et al. Triptan-induced contractile (5-HT_{1B} receptor) responses in human cerebral and coronary arteries: relationship to clinical effect. *Clin Sci (Lond)*. 2005; 109:335-342 _____ regel 7
229. Humphrey PP, Goadsby PJ. The mode of action of sumatriptan is vascular? A debate. *Cephalalgia*. 1994; 14:401-410 _____ regel 8
230. Villalon CM, Centurion D, Valdivia LF et al. Migraine: pathophysiology, pharmacology, treatment and future trends. *Curr Vasc Pharmacol*. 2003; 1:71-84 _____ regel 9
231. Hoskin KL, Kaube H, Goadsby PJ. Sumatriptan can inhibit trigeminal afferents by an exclusively neural mechanism. *Brain*. 1996; 119:1419-1428 _____ regel 10
232. Hoskin KL, Kaube H, Goadsby PJ. Central activation of the trigeminovascular pathway in the cat is inhibited by dihydroergotamine. A c-Fos and electrophysiological study. *Brain*. 1996; 119 (Pt 1):249-256 _____ regel 11
233. Goadsby PJ. Can we develop neurally acting drugs for the treatment of migraine? *Nat Rev Drug Discov*. 2005; 4:741-750 _____ regel 12
234. Ferrari MD, Roon KI, Lipton RB et al. Oral triptans (serotonin 5-HT_{1B/1D} agonists) in acute migraine treatment: a meta-analysis of 53 trials. *Lancet*. 2001; 358:1668-1675 _____ regel 13
235. Dodick D, Lipton RB, Martin V et al. Consensus statement: cardiovascular safety profile of triptans (5-HT agonists) in the acute treatment of migraine. *Headache*. 2004; 44:414-425 _____ regel 14
236. Masuzawa T, Shinoda S, Furuse M et al. Cerebral angiographic changes on serial examination of a patient with migraine. *Neuroradiology*. 1983; 24:277-281 _____ regel 15
237. Markus HS. Transcranial Doppler ultrasound. *Br Med Bull*. 2000; 56:378-388 _____ regel 16
238. Krabbe-Hartkamp MJ, Grond van der J, de Leeuw F-E et al. Circle of Willis: Morphologic variation on three-dimensional time-of-flight MR angiograms. *Radiology*. 1998; 207:103-111 _____ regel 17
239. Iversen HK, Olesen J. Headache induced by a nitric oxide donor (nitroglycerin) responds to sumatriptan. A human model for development of migraine drugs. *Cephalalgia*. 1996; 16:412-418 _____ regel 18

References

- regel 1 _____ 240. Andresen J, Shafi NI, Bryan RM, Jr. Endothelial influences on cerebrovascular tone. *J Appl Physiol*. 2006; 100:318-327
- regel 2 _____
- regel 3 _____ 241. Strecker T, Dux M, Messlinger K. Increase in meningeal blood flow by nitric oxide--interaction with calcitonin gene-related peptide receptor and prostaglandin synthesis inhibition. *Cephalalgia*. 2002; 22:233-241
- regel 4 _____
- regel 5 _____
- regel 6 _____ 242. Abrams J. Pharmacology of nitroglycerin and long-acting nitrates. *Am J Cardiol*. 1985; 56:12A-18A
- regel 7 _____ 243. Caekebeke JF, Ferrari MD, Zwetsloot CP et al. Antimigraine drug sumatriptan increases blood flow velocity in large cerebral arteries during migraine attacks. *Neurology*. 1992; 42:1522-1526
- regel 8 _____
- regel 9 _____ 244. Limmroth V, May A, Auerbach P et al. Changes in cerebral blood flow velocity after treatment with sumatriptan or placebo and implications for the pathophysiology of migraine. *J Neurol Sci*. 1996; 138:60-65
- regel 10 _____
- regel 11 _____
- regel 12 _____ 245. Gori S, Morelli N, Bellini G et al. Rizatriptan does not change cerebral blood flow velocity during migraine attacks. *Brain Res Bull*. 2005; 65:297-300
- regel 13 _____
- regel 14 _____ 246. Zwetsloot CP, Caekebeke JFV, Ferrari MD. Lack of asymmetry of middle cerebral artery blood velocity in unilateral migraine. *Stroke*. 1993; 24:1335-1338
- regel 15 _____
- regel 16 _____ 247. Guyton AC. Cerebral Blood Flow, Cerebrospinal Fluid and Brain Metabolism. In: Guyton AC, Hall JE, eds. *Textbook of medical physiology*. Philadelphia: Saunders, 2006:761-768
- regel 17 _____
- regel 18 _____ 248. Afridi S, Kaube H, Goadsby PJ. Occipital activation in glyceryl trinitrate induced migraine with visual aura. *J Neurol Neurosurg Psychiatry*. 2005; 76:1158-1160
- regel 19 _____
- regel 20 _____ 249. Kruit MC, Launer LJ, Ferrari MD et al. Brain stem and cerebellar hyperintense lesions in migraine. *Stroke*. 2006; 37:1109-1112
- regel 21 _____
- regel 22 _____ 250. Kruit MC, van Buchem MA, Hofman PA et al. Migraine as a risk factor for subclinical brain lesions. *JAMA*. 2004; 291:427-434
- regel 23 _____
- regel 24 _____ 251. de Leeuw R, Schmidt JE, Carlson CR. Traumatic stressors and post-traumatic stress disorder symptoms in headache patients. *Headache*. 2005; 45:1365-1374
- regel 25 _____
- regel 26 _____ 252. Christiansen I, Daugaard D, Lykke TL et al. Glyceryl trinitrate induced headache in migraineurs - relation to attack frequency. *Eur J Neurol*. 2000; 7:405-411
- regel 27 _____
- regel 28 _____ 253. Vanmolkot KR, Kors EE, Hottenga JJ et al. Novel mutations in the Na⁺, K⁺-ATPase pump gene ATP1A2 associated with familial hemiplegic migraine and benign familial infantile convulsions. *Ann Neurol*. 2003; 54:360-366
- regel 29 _____
- regel 30 _____
- regel 31 _____ 254. Dreier JP, Jurkat-Rott K, Petzold GC et al. Opening of the blood-brain barrier preceding cortical edema in a severe attack of FHM type II. *Neurology*. 2005; 64:2145-2147
- regel 32 _____
- regel 33 _____ 255. Olesen J, Thomsen LL, Iversen HK. Nitric oxide is a key molecule in migraine and other vascular headaches. *Trends Pharmacol Sci*. 1995; 15:149-153
- regel 34 _____
- regel 35 _____
- regel 36 _____
- regel 37 _____
- regel 38 _____
- regel 39 _____

BIBLIOGRAPHY

N Schmitz, EB Arkink, M Mulder, K Rubia, F Admiraal-Behloul, GG Schoonman, MC Kruit, MD Ferrari, MA van Buchem. Frontal lobe structure and executive function in migraine patients. *Neurosci Lett*. 2008 May 16. (Epub ahead of print)

GG Schoonman, J van der Grond, GM Terwindt, MD Ferrari. Cerebral blood flow response to nitroglycerin predicts the occurrence of a provoked migraine attack. Submitted

GG Schoonman, J van der Grond, C Kortman, RJ van der Geest, GM Terwindt, MD Ferrari. Migraine headache is not associated with cerebral or meningeal vasodilatation - a 3T magnetic resonance angiography study. *Brain*. 2008 May 23 (Epub ahead of print)

N Schmitz, F Admiraal-Behloul, EB Arkink, MC Kruit, GG Schoonman, MD Ferrari, MA van Buchem. Attack Frequency and Disease Duration as Indicators for Brain Damage in Migraine. *Headache*. 2008 May 9. (Epub ahead of print)

D Magis, L Bendtsen, PJ Goadsby, A May, M Sánchez del Rio, PS Sandór, H Kaube, G Sandrini, GG Schoonman, J Schoenen. Evaluation and proposal for optimisation of neurophysiological tests in migraine: Neuroimaging and the nitroglycerin test. *Cephalalgia*. 2007;27:1339-59.

GG Schoonman, PS Sándor, AC Nirkko, T Lange, T Jaermann, U Dydak, C Kremer, MD Ferrari, P Boesiger, RW Baumgartner. Experimental hypoxia induced acute mountain sickness is associated with intracellular cerebral oedema. A 3 Tesla Magnetic resonance imaging study. *Journal of Cerebral Blood Flow and Metabolism*. 2008; 28:198-206.

GG Schoonman, DJ Evers, BE Ballieux, EJ de Geus, ER de Kloet, GM Terwindt, JG van Dijk, MD Ferrari. Is stress a trigger factor for migraine? *Psychoneuroendocrinology* 2007;32: 532-538.

GG Schoonman, D Bakker, N Schmitz, RJ van der Geest, J van der Grond, MD Ferrari, MA van Buchem. Magnetic Resonance Angiography of the Human Middle Meningeal Artery: Implications for Migraine. *J Magn Reson Imaging*. 2006; 24: 918-21.

___ regel 1

___ regel 2

___ regel 3

___ regel 4

___ regel 5

___ regel 6

___ regel 7

___ regel 8

___ regel 9

___ regel 10

___ regel 11

___ regel 12

___ regel 13

___ regel 14

___ regel 15

___ regel 16

___ regel 17

___ regel 18

___ regel 19

___ regel 20

___ regel 21

___ regel 22

___ regel 23

___ regel 24

___ regel 25

___ regel 26

___ regel 27

___ regel 28

___ regel 29

___ regel 30

___ regel 31

___ regel 32

___ regel 33

___ regel 34

___ regel 35

___ regel 36

___ regel 37

___ regel 38

___ regel 39

Bibliography

- regel 1 _____ GG Schoonman, DJ Evers, GM Terwindt, JG van Dijk, MD Ferrari. The prevalence of
regel 2 _____ premonitory symptoms in migraine: a questionnaire study in 461 patients. Cephalalgia
regel 3 _____ 2006; 26: 1209-13.
- regel 4 _____
- regel 5 _____ GG Schoonman, PS Sándor, RM Agosti, M Siccoli, P Bärtsch, MD Ferrari, RW Baumgartner.
regel 6 _____ Normobaric hypoxia and nitroglycerin as trigger factors for migraine. Cephalalgia 2006;
regel 7 _____ 26: 816-9.
- regel 8 _____
- regel 9 _____ GG Schoonman, NJ Wiendels, MD Ferrari. Acupunctuur bij de profylactische behandeling
regel 10 _____ van migraine; voorlopig geen bewijs van effectiviteit. Ned Tijdschr Geneeskunde
regel 11 _____ 2004;148: 2165-6
- regel 12 _____
- regel 13 _____ GG Schoonman, NJ Wiendels, MD Ferrari. Gabapentin in migraine prophylaxis: is it
regel 14 _____ effective and well tolerated? Headache 2002; 42: 235
- regel 15 _____
- regel 16 _____ KI Roon, PS Sándor, GG Schoonman, FPL Lamers, J Schoenen, MD Ferrari, JG van Dijk.
regel 17 _____ Auditory evoked potentials in the assessment of central nervous system effects of
regel 18 _____ antimigraine drugs. Cephalalgia 1999; 19: 880-5
- regel 19 _____
- regel 20 _____ J Haan, GG Schoonman, GM Terwindt, MD Ferrari, RM Bertina, LJ Kappelle.
regel 21 _____ Prothrombotic mutations and ischaemic stroke at a young age in two sisters. J Neurol
regel 22 _____ Neurosurg Psychiatry 1998; 65: 958-9
- regel 23 _____
- regel 24 _____
- regel 25 _____
- regel 26 _____
- regel 27 _____
- regel 28 _____
- regel 29 _____
- regel 30 _____
- regel 31 _____
- regel 32 _____
- regel 33 _____
- regel 34 _____
- regel 35 _____
- regel 36 _____
- regel 37 _____
- regel 38 _____
- regel 39 _____

CURRICULUM VITAE

Geurt Gerhard Schoonman (roepnaam: Guus) werd geboren op 14 oktober 1974 in Deventer. Hij behaalde in 1993 zijn VWO diploma aan het Baudartius College te Zuthpen. Het doctoraalexamen Geneeskunde behaalde hij in 1998 aan de Universiteit Leiden. De wetenschappelijke stage (titel: "Decompression sickness and pyramidal tract demyelination in rats") werd gelopen aan de Universiteit van Kopenhagen (supervisor: dr. M. Ballegaard). Daarnaast heeft hij tussen 1994 en 1998 psychologie gestudeerd aan de Universiteit Leiden. De co-schappen in het Academisch Ziekenhuis Leiden (AZL, nu Leids Universitair Medisch Centrum, LUMC) sloot hij af met het artsexamen in 2000. Gedurende 6 jaar was hij arts-onderzoeker in het LUMC onder begeleiding van Prof. Dr. M.D. Ferrari. In 2002 was hij 6 maanden in Zwitserland voor wetenschappelijk onderzoek naar de relatie tussen hypoxie en migraine aan de Universiteit van Zurich (supervisor: Prof. Dr. R.W. Baumgartner). Naast wetenschappelijk onderzoeker was hij voorzitter van de Trainees and Residents subcommittee van de International Headache Society en bestuurslid van de vereniging voor arts-onderzoekers in het LUMC. In 2007 startte hij als arts-assistent neurologie in het Diaconnessenhuis te Leiden. Momenteel is hij wederom werkzaam in het LUMC als arts-assistent in opleiding tot neuroloog (opleider: Prof. Dr. R.A.C. Roos).

— regel 1
— regel 2
— regel 3
— regel 4
— regel 5
— regel 6
— regel 7
— regel 8
— regel 9
— regel 10
— regel 11
— regel 12
— regel 13
— regel 14
— regel 15
— regel 16
— regel 17
— regel 18
— regel 19
— regel 20
— regel 21
— regel 22
— regel 23
— regel 24
— regel 25
— regel 26
— regel 27
— regel 28
— regel 29
— regel 30
— regel 31
— regel 32
— regel 33
— regel 34
— regel 35
— regel 36
— regel 37
— regel 38
— regel 39

— |

| —

— |

| —