Influence of Acute Jugular Vein Compression on the Cerebral Blood Flow Velocity, Pial Artery Pulsation and Width of Subarachnoid Space in Humans

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Abstract

Purpose: The aim of this study was to assess the effect of acute bilateral jugular vein compression on: (1) pial artery pulsation (cc-TQ); (2) cerebral blood flow velocity (CBFV); (3) peripheral blood pressure; and (4) possible relations between mentioned parameters.

Methods: Experiments were performed on a group of 32 healthy 19–30 years old male subjects. cc-TQ and the subarachnoid width (sas-TQ) were measured using near-infrared transillumination/backscattering sounding (NIR-T/BSS), CBFV in the left anterior cerebral artery using transcranial Doppler, blood pressure was measured using Finapres, while end-tidal CO₂ was measured using medical gas analyser. Bilateral jugular vein compression was achieved with the use of a sphygmomanometer held on the neck of the participant and pumped at the pressure of 40 mmHg, and was performed in the bend-over (BOPT) and swayed to the back (initial) position.

Results: In the first group (n = 10) during BOPT, sas-TQ and pulse pressure (PP) decreased (−17.6% and −17.9%, respectively) and CBFV increased (+35.0%), while cc-TQ did not change (+1.9%). In the second group, in the initial position (n = 22) cc-TQ and CBFV increased (106.6% and 20.1%, respectively), while sas-TQ and PP decreases were not statistically significant (−15.5% and −9.0%, respectively). End-tidal CO₂ remained stable during BOPT and venous compression in both groups. Significant interdependence between changes in cc-TQ and PP after bilateral jugular vein compression in the initial position was found (r = −0.74).

Conclusions: Acute bilateral jugular venous insufficiency leads to hyperkinetic cerebral circulation characterised by augmented pial artery pulsation and CBFV and direct transmission of PP into the brain microcirculation. The Windkessel effect with impaired jugular outflow and more likely increased intracranial pressure is described. This study clarifies the potential mechanism linking jugular outflow insufficiency with arterial small vessel cerebral disease.

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Competing Interests: This study was partially funded by NIRT sp. z o.o., Wierzbie, Poland, who also rented the authors the SAS-Monitor device, which is a product in development. A. Frydrychowski owns several patents related to NIR-T/BSS technology and is a stakeholder in NIRT sp. z o.o. Patents: A. Frydrychowski, M. Rukasz: P.376246, 28 November 2011; A. Frydrychowski, M. Rukasz: P.357040, 21 March 2007; A. Frydrychowski, M. Rukasz: P.344496, 31 July 2007. All patents are related to non-invasive monitoring of the subarachnoid space and pial artery, signal detection and analysis. There are no further patents, products in development or marketed products to declare. This does not alter the authors’ adherence to all the PLOS ONE policies on sharing data and materials.

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Introduction

A stable increase in cerebral venous pressure induces arteriolar vasodilation, which averages 6% to 12% of the control diameter in cats [1,2] and dogs [3]. Acute superior vena cava occlusion results in increased pial venous pressure and subsequent blood-brain barrier disruption in rats [4]. Unilateral internal and external jugular vein ligation in swine increases bihemispheric cerebral blood flow and metabolism [5], and progressive superior vena cava occlusion produces measurable signs of impaired cerebral perfusion in pigs [6]. As cranial window installation and microscopic examination remain the methods of choice to investigate the pial microvessels in vivo [7], for obvious reasons, the effect of increased cerebral venous pressure has not yet been investigated in humans.

The amplitude of pial artery pulsation (cc-TQ) can be measured non-invasively using near-infrared transillumination/backscattering sounding (NIR-T/BSS), a new method based on infrared radiation (IR) that has been developed in the last decade by our team [8–12]. Contrary to near-infrared spectroscopy (NIRS), which relies on absorption of IR by haemoglobin [13,14], NIR-T/BSS uses the subarachnoid space (SAS) filled with translucent
cerebrospinal fluid (CSF) as a propagation duct for IR. We have previously shown that pial artery pulsation may serve as a sensitive index of changes in microvessel compliance. cc-TQ increases were observed during acetazolamide and hypercapnic tests, acute hypoxia, papaverine and glucose administration and electroconvulsive therapy, while cc-TQ decreases were recorded during the stabilisation period after the abovementioned procedures [10,15–17]. Significant cc-TQ decrease was seen during handgrip test [18]. In addition, NIR-T/BSS allows for assessment of changes in the width of SAS [19], indicative in changes in CSF volume and intracranial pressure [8,10–12,20]. Due to non-invasiveness, ease of use and low cost NIR-T/BSS potentially constitutes an ideal tool to monitor brain microcirculation over long periods of time.

We hypothesised that impaired venous outflow would lead to pial artery dilation caused by a metabolic and/or autoregulatory mechanism due to decreased oxygen supply and/or increased intracranial pressure, respectively. Augmented pial artery compliance should result in increased cc-TQ and hyperdynamic brain circulation characterised by augmented cerebral blood flow velocity (CBFV). The investigated topic is not a purely theoretical concept, as increased cerebral venous blood volume has been recently reported during sympathetic activation (for a review, see [21]). Therefore, the brain may be exposed to venous congestion much more frequently than initially thought. Furthermore, venous dysfunction is increasingly linked to pulse wave encephalopathy and white matter changes [22–25]. The aim of this study was to assess in healthy volunteers the effect of acute jugular venous compression on: (1) cc-TQ; (2) CBFV; (3) peripheral blood pressure (BP); and possible relations between CBFV, BP and cc-TQ.

Materials and Methods

Subjects

Experiments were performed on a group of 32 healthy 19–30 year old male subjects. The volunteers were selected on the basis of a medical questionnaire, interview and blood pressure measurement. All volunteers received detailed information about study objectives and potential adverse reactions (transient headache, vertigo and blood flushes) and gave written informed consent to participate in the study. The experimental protocol and the study were approved by the ethical committee of the Medical University of Gdańsk (TKEBN/259). Subjects did not have any disorders and were not taking any medication. Before the experiment, each subject underwent a general and neurological examination. No nicotine, coffee, tea, cocoa or any methylnitrothine-containing food or beverages were permitted for 8 hours before the tests. Additionally, prior to each test, the volunteers were asked to sit comfortably and rest for 30 minutes.

Experimental design

**Bend Over Position Test (BOPT)**

- Initial position: subject set comfortably for 10 minutes with head swayed to the back at 20° angle.
- BOPT: subject bends forwards at 45° angle (the trunk and the head) from the initial half-lying position on the back [10,18].

Then, for bilateral jugular vein compression participants were divided into two groups.

**First group consisting of 10 subjects**

- Jugular veins were completely compressed 3 minutes after starting BOPT for 3 minutes.
- Subject returned to initial position.

**Second group consisting of 22 subjects**

- Subject returned to initial position 3 minutes after starting BOPT.
- Subject set comfortably in initial position for 3 minutes.
- Jugular veins were completely compressed for 3 minutes.

The numerical values for statistical analysis were taken just before the end of each three-minute procedure. After bilateral jugular vein compression all subject were asked to sit comfortably in the initial position for 10 minutes.

Bilateral jugular vein compression was achieved with the use of a sphygmomanometer held on the neck of the participant and pumped to a pressure of 40 mmHg. Blood pressure in jugular veins in the initial position should be close to zero, or even negative, so it can be assumed that compression was complete. The subjects were asked to breathe normally to avoid changes in PaCO2.

**Transcranial Doppler measurement**

Transcranial Doppler (TCD) measurement of cerebral blood flow velocity (CBFV) in the left anterior cerebral artery was performed with a Doppler ultrasound device TDS 4 (Sonomed, Warsaw, Poland). A pulse probe of 2 MHz was used and analysis of the results was carried out on a built-in IBM PC computer. To assure the best possible reproducibility of the recordings, the Doppler probe was mounted on the head with a special stabilising strip.

**Blood pressure measurement**

Recording of changes in systolic arterial pressure (SAP), diastolic arterial pressure (DAP) and HR were measured using Finapres (Finapres, Ohmmeda, Englewood, CO, USA). The Finapres sensor was mounted onto the middle finger of the non-dominant hand resting on the table. Beat-to-beat blood pressure was transferred to a computer console continuously displaying SAP, DAP and HR.

Pulse pressure (PP) was calculated from the following equation: $PP = SAP - DAP$.

**End-tidal CO2 measurement**

End-tidal CO2 was measured in mmHg with a medical gas analyser (Datex Instrumentarium, Helsinki, Finland). The instrument was calibrated using a certified standard gas mixture before each experiment. End-tidal CO2 was recorded and printed continuously in real time. Values for subsequent analysis were read from the printouts and entered manually into a Statistics for Windows 8.0 database.

**NIR-T/BSS measurement**

Recording of changes in the amplitude of pial artery pulsation and in the width of SAS with NIR-T/BSS were performed with a head-mounted SAS 100 Monitor (NIRT sp. z o.o., Wierzbice, Poland). The sensor unit consists of the emitter (E) and two photosensors located at different distances from the emitter. The NIR-T/BSS emitter is a near-infrared light-emitting diode (LED). The proximal sensor (PS) is located close to the emitter, while the distal sensor (DS) is located at a larger distance from the emitter. The stream of IR generated by the emitter penetrates the highly perfuse layer of the skin of the head, the skull bones and the SAS. The stream of radiation reflects from the surface of the brain and reaches the sensors, crossing the aforementioned layers of tissues in reverse order. Signals from the sensors undergo analogue-digital conversion in a specialised data acquisition system and are recorded on the microcomputer’s hard disk for subsequent analysis with on-line presentation on the computer monitor.

Theoretical and practical foundations of the NIR-T/BSS method were provided in the model studies by our team [8,10–
Briefly, a signal received by the DS is divided by the signal received by the PS. Such division reduces proportional factors that affect each of the two signals in an identical way, because the quotient of these factors assumes the value 1. Both the dividend, i.e., the power of the DS signal, and the divisor, i.e., the power of the PS signal, are influenced by the width of SAS, and also by any factor capable of changing that width. Therefore, the quotient of the two signals, called the transillumination quotient (TQ), is

Figure 1. Effect of acute bilateral jugular vein compression on NIR-T/BSS variables during BOPT: cc-TQ is “cut” by the narrowing SAS. Sharp edges of the cc-TQ waves are visible at the distant sensor (DS) and cc-TQ (enhanced tracings). cc-TQ – cardiac component of transillumination quotient (pial artery pulsation); μW/cm² – microwatt/centimetre².

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sensitive to changes in the width of the SAS. The oscillations of TQ have their origin in different modulation of the PS and DS signals, namely in the modulation of the DS signal on its way through the SAS. It happens because only the DS receives radiation propagated within the SAS. Propagation of IR in the skin and bone is much worse than in the clear translucent cerebrospinal fluid (CSF) contained in the SAS, and with the DS placed far enough from the emitter, no radiation propagated in the superficial tissue layers can reach the DS [8,10–12,20]. The power of the IR stream reaching the DS is directly proportional to the width of the SAS. The wider the SAS, or the propagation duct, the more radiation reaches the DS and the greater the signal from that sensor, which is the dividend in the calculation of the TQ [8,10–12,20]. Conversely, the power of the IR stream reaching the PS is inversely proportional to the width of the SAS. Thus, the wider the SAS, the longer the distance between the inner skull bone surface and the brain at which IR energy is dissipated, and therefore the less IR follows the return route to the PS. The narrower the SAS, the closer the reflecting surface of the brain is to the PS and the more IR reaches that sensor.

Thus, in the transillumination quotient (TQ) we can identify three main components:

1. constant or a non-pulsatile component, further referred to as the subarachnoid component (sas-TQ); its value depending on the width of the CSF-filled SAS,
2. slow-variable pulsation, further referred to as the subcardiac component (scc-TQ); including, but not limited to, pulsation of a respiratory origin,
3. fast-variable pulsation, further referred to as the cardiac component (cc-TQ); resulting from heart-generated arterial pulsation that is the cause of fast oscillations of the width of SAS.

For further analysis, the first harmonic of the arterial pulsation-dependent oscillations of TQ was extracted through appropriate filtering, along with its modulation. Modulation of that harmonic is a fast-variable component (or cardiac component) of principal, second, and third harmonics of cardiac component waveform, respectively.

Statistical analysis

W Shapiro-Wilk W, Mann-Whitney U and ANOVA tests were used to analyse the differences between average values. Changes in sas-TQ, cc-TQ, CBFV, SAP, DAP, PP, HR and end-tidal CO2 responses were compared to baseline values. Correlation and regression analysis was performed to assess interdependences between SAP, DAP, PP, HR, CBFV, end-tidal CO2, sas-TQ and cc-TQ. All statistical calculations were performed using the Statistics for Windows 8.0 commercial package.

Results

Both groups (acute jugular vein compression in BOPT and initial position) were well-matched at baseline in terms of CBFV and BP. The BOPT test caused sas-TQ, cc-TQ, CBFV and PP decrease in both investigated groups (n = 10 and n = 22). However, acute bilateral jugular vein compression brought different results between the investigated groups. In the first group (n = 10) during
BOPT sas-TQ and PP further decreased, CBFV increased while cc-TQ did not change (Fig. 1). In the second group, after return to initial position, during bilateral jugular vein compression (n = 22) cc-TQ and CBFV increased, while sas-TQ and PP decrease were not statistically significant (Fig. 2). End-tidal CO2 remained stable during BOPT and venous compression in both groups. Detailed descriptive statistics are provided in Table 1 and Table 2. Changes in end-tidal CO2 were not clinically relevant. Therefore, venous congestion is independent of hypercapnia in the induction of increased cerebral blood flow. Thus, the current study contributes to a better understanding of the mechanisms underlying the pathology of diseases characterised by increased cerebral venous blood volume.

The brain is subject to gravitation and changes its position along with changes in the position of the head. At a forward bending position (during BOPT) or in the abdominal-lying position, the brain floats forward toward the inner surface of the frontal bones for the first time, confirming the results of earlier animal studies [1–3].

All changes are calculated versus preceding values. *p<0.05; **p<0.01; ***p<0.001. sas-TQ – the subarachnoid component of the transillumination quotient (the subarachnoid width); cc-TQ – cardiac component of transillumination quotient (pial artery pulsation); CBFV – cerebral blood flow velocity; SAP – systolic arterial pressure; DAP – diastolic arterial pressure; PP – pulse pressure; HR – heart rate.

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### Discussion

This study, to the best of our knowledge, describes the effects of venous congestion on the pial arteries and CBFV in humans for the first time, confirming the results of earlier animal studies [1–3]. The new findings are that acute bilateral jugular outflow congestion in humans leads to hyperkinetic cerebral circulation, increased pial artery pulsation and direct transmission of peripheral PP into pial arteries. It is unlikely that the hyperdynamic brain circulation was caused by increased PaCO2 since the changes in end-tidal CO2 were not clinically relevant. Therefore, venous congestion is independent of hypercapnia in the induction of increased cerebral blood flow. Thus, the current study contributes to a better understanding of the mechanisms underlying the pathology of diseases characterised by increased cerebral venous blood volume.

The brain is subject to gravitation and changes its position along with changes in the position of the head. At a forward bending position (during BOPT) or in the abdominal-lying position, the brain floats forward toward the inner surface of the frontal bones for the first time, confirming the results of earlier animal studies [1–3].

### Table 1. Effects of acute bilateral jugular vein compression on sas-TQ, cc-TQ, CBFV, SAP, DAP, PP and HR during the BOPT test (BOPT – JVO; n = 10). Mean values and Standard Deviations (SD) are provided.

|          | Initial position (baseline) | BOPT% Change | BOPT – JVO% Change | Initial position/recovery % Change |
|----------|----------------------------|--------------|--------------------|-----------------------------------|
| sas-TQ (arbitrary units) | 1623±55.7 | 1123±48.1*** | −30.8 | 926±36.7*** | −17.6 | 1178±50.8** | 27.2 |
| cc-TQ (arbitrary units) | 54.5±24.2 | 34.0±12.8** | −37.5 | 34.7±17.0NS | 1.9 | 33.3±11.3** | −3.95 |
| CBFV (cm\*sec⁻¹) | 59.3±14.1 | 45.1±9.7** | −24.0 | 60.9±11.6** | 35.0 | 51.8±13.4** | −15.0 |
| SAP (mmHg) | 138.0±12.3 | 131.0±12.0** | −5.07 | 131.7±17.3NS | 0.53 | 126.5±14.3** | −3.95 |
| DAP (mmHg) | 74.1±9.1 | 73.9±9.2NS | −0.27 | 84.8±12.4** | 14.75 | 73.6±9.7NS | −13.2 |
| PP (mmHg) | 63.9±8.8 | 57.1±7.9** | −10.6 | 46.9±14.2** | −17.9 | 52.9±12.1** | 12.8 |
| HR (beats*sec⁻¹) | 75.6±7.9 | 78.3±6.4NS | 3.57 | 83.5±9.6* | 6.64 | 52.9±12.8** | −6.83 |
| End-tidal CO2 (mmHg) | 35.6±1.7 | 36.8±3.2NS | 3.37 | 36.3±2.9NS | −1.4 | 36.1±5.3NS | −0.56 |

### Table 2. Effects of acute bilateral jugular vein compression on sas-TQ, cc-TQ, CBFV, SAP, DAP, PP and HR in initial position (Initial – JVO; n = 22).

|          | Initial position (baseline) | BOPT% Change | Initial position/recovery after BOPT% Change | Initial – JVO % Change |
|----------|----------------------------|--------------|---------------------------------------------|------------------------|
| sas-TQ (arbitrary units) | 1845±81.2 | 1287±61.8*** | −30.2 | 1861±78.5*** | 44.6 | 1573±79.8NS | −15.5 |
| cc-TQ (arbitrary units) | 70.8±53.7 | 42.8±41.6* | −39.7 | 67.1±54.4* | 56.8 | 138.6±32.8* | 106.6 |
| CBFV (cm\*sec⁻¹) | 62.3±11.7 | 47.8±10.4** | −23.3 | 61.2±17.4** | 28.0 | 73.5±15.4** | 20.1 |
| SAP (mmHg) | 140.8±11.9 | 131.5±18.2** | −6.7 | 140.6±12.0NS | 6.9 | 140.3±10.5 NS | −0.03 |
| DAP (mmHg) | 79.7±9.2 | 78.3±9.4NS | −1.7 | 79.4±8.7* | 1.4 | 82.6±7.9 NS | 4.0 |
| PP (mmHg) | 61.1±7.0 | 53.1±15.5* | −13.1 | 61.2±7.1** | 15.3 | 57.7±5.0NS | −9.0 |
| HR (beats*sec⁻¹) | 77.9±10.4 | 79.0±11.3NS | 1.3 | 77.4±9.9NS | 2.0 | 79.0±10.0* | 2.1 |
| End-tidal CO2 (mmHg) | 37.2±4.3 | 36.4±3.1NS | −2.2 | 36.8±3.6NS | 1.1 | 37.1±4.8NS | −0.8 |

Mean values and Standard Deviations (SD) are provided. All changes are calculated versus preceding values. *p<0.05; **p<0.01; ***p<0.001. sas-TQ – the subarachnoid component of the transillumination quotient (the subarachnoid width); cc-TQ – cardiac component of transillumination quotient (pial artery pulsation); CBFV – cerebral blood flow velocity; SAP – systolic arterial pressure; DAP – diastolic arterial pressure; PP – pulse pressure; HR – heart rate.

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and the SAS in the frontal region assumes its minimum value [19]. The observed decreases in sas-TQ and cc-TQ during the BOPT have already been discussed in detail [10,18,19]. What is important, and has not been presented earlier, is that decreases in cc-TQ are fully consistent with CBFV decreases observed in TCD. Hayreh and Edwards [26] demonstrated that during the very early phase of intracranial pressure elevation PP may decrease. Importantly, Wein and Kontos [2] reported a small but consistent decrease in BP after the onset of venous hypertension. Thus, the observed decrease in PP may suggest a slight increase in intracranial pressure during BOPT. However, based on the NIR-T/BSS model, we are not able to distinguish if a decrease in sas-TQ is solely due to physical brain movement or is in addition influenced by increased intracranial pressure.

Bilateral jugular vein compression resulted in enhanced CBFV in both groups. We may assume that hampered jugular outflow results in decreased oxygen supply, which in turn lead to arteriolar dilation and subsequent increase in cerebral blood flow [5]. Pial artery dilation and increased compliance results in higher amplitude of pial artery pulsation [10,15–17]. Alternatively, bilateral jugular vein compression more likely evoked an increase in intracranial pressure. Even a mild increase in intracranial pressure results in an autoregulatory response to diminished cerebral perfusion pressure, i.e. pial artery dilation and increased pulsation [10,12,20]. Therefore, cc-TQ should have been increased too. However, when jugular veins were compressed during BOPT, cc-TQ did not change. To explain this result we need to focus on assumptions built in the NIR-T/BSS model. NIR-T/BSS uses the SAS filled with CSF as a propagation duct for IR. Therefore the presence of SAS is the sine qua non condition for measurements of any NIR-T/BSS parameters. The SAS already decreased during BOPT gets even more squeezed during acute bilateral jugular vein compression, due to expanded intracranial blood volume, and more likely increased intracranial pressure. Therefore, the “cutting” effect actually develops due to the Windkessel mechanism. In order to compensate for higher resistance in the venous drainage pathways, the pial artery tries to dilate. However, this is not possible because the SAS is compressed and non-compliant. What we can see in Fig. 1 is that the amplitude of CVP is being cut by the narrowing SAS. This is an extremely important feature of NIR-T/BSS as the “cutting” of pial artery pulsation can be indicative of early brain oedema in human (unpublished results from our lab). BOPT, which was developed by our team 10 years ago, can be used to non-invasively assess brain reserve volume. Nevertheless, the above effect does not allow for assessment of acute bilateral vein compression influence on cc-TQ during BOPT. The “cutting” effect was observed in all volunteers during BOPT (n = 10). As the aim of this study was to assess pial artery pulsation during acute bilateral vein compression, we decided not to enrol more subjects into this experimental group.

Acute cerebral venous congestion produced significant cc-TQ increase in the initial position. Such a result is in line with the above-presented reasoning. In the initial position the SAS was insignificantly reduced by enhanced intracraniac blood volume, and remained wide enough to allow for proper registration of the amplitude of pial artery pulsation (Fig. 2). Increases in CBFV and pial artery pulsation observed during acute impairment of venous outflow are consistent with the model proposed by Bateman [25]. Pulsatile flow is a manifestation of the energy stored in the form of the pulse pressure and, to ensure non-pulsatile continuous flow through the capillary bed, must be dampened by shifting cerebrospinal fluid and venous blood [27]. When venous outflow is impaired and intracranial pressure is elevated, the pulse pressure cannot be dampened and results in increased pial artery pulsation [28]. Correlation and regression analysis revealed strong interdependence between cc-TQ and PP during acute bilateral jugular vein compression in the initial position (Fig. 3). Exposure to highly pulsatile pressure and augmented flow is a known predictor of cerebral vascular damage, even in the absence of increases in mean BP [29–31]. We have already proved in an animal model that during hypercapnia pulsatile flow is directly transmitted into pial arteries [16]. The presented correlation may actually provide a link between cerebral venous insufficiency and cerebral arterial small vessel disease in human. Chronic cerebral insufficiency is associated with impaired cerebral perfusion [6,32]. Toveda et al. [6] and Zamboni et al. [32] reports are not contradictory to our results. The Windkessel mechanism requires the SAS to be compliant to ensure smooth blood flow through the cerebral vascular bed. If the pial arteries cannot expand, then the flow through the vascular bed becomes more pulsatile and the shear forces on the arterial wall increase [23]. Direct transmission of PP into the brain microcirculation, if maintained over a longer period of time, may result in subsequent arteriole remodelling, micro-bleeding and increased vascular resistance, leading to an augmented vulnerability to ischaemia as the final outcome [29–31]. Furthermore, elevated arterial inflow combined with impaired venous outflow may lead to the development of normal pressure hydrocephalus [23], a disease associated with leukoaraiosis [22,24,25], altered cerebral pulsation propagation [22] and reduced cerebral blood flow [33]. Zamboni reported [32] a link between chronic cerebrospinal venous insufficiency and hypoperfusion in multiple sclerosis; although this remains controversial, it is in line with the reasoning presented above.

The following study limitations should be taken into account. Only young, healthy males were investigated and acute jugular vein insufficiency was analysed. Therefore, any extrapolations to patients suffering from chronic cerebral venous insufficiency or normal pressure hydrocephalus should be viewed with caution. In our study the SAS decrease during bilateral jugular vein compression in the initial position was not statistically significant, which may suggest that even during bilateral jugular vein congestion the brain may be decompressed through the vertebral plexus [34] or venous collateral circulation [35]. The high within-
and between-subject reproducibility and repeatability of NIR-T/BSS measurements have been demonstrated earlier [9,10,19,36]. NIR-T/BSS, like NIRS, allows for direct within-subject comparisons. However, it should not be noted that the percentage changes during BOPT were almost identical in both groups. This is in agreement with our previous studies showing that as long as changes from baseline values are analysed, high between-subject reproducibility is observed [10,16–19]. Studies showing that as long as changes from baseline values are related to differences in skull bone parameters. However, it should actually exaggerates the pulsatile flow, which in turn creates a hazardous environment for the brain microcirculation. Therefore, this study clarifies the potential mechanism linking jugular outflow insufficiency with small vessel arterial cerebral disease.

**References**

1. Wei EP, Kontos HA (1982) Responses of cerebral arterioles to increased venous pressure. Am J Physiol 243: H447–9.

2. Wei EP, Kontos HA (1984) Increased venous pressure causes myogenic constriction of cerebral arterioles during local hyperxia. Circ Res 55:249–52.

3. Wagner EM, Traystman RJ (1983) Cerebral venous outflow and arterial microsphere flow with elevated venous pressure. Am J Physiol 244: H505–12.

4. Mayhan WG, Heistad DD (1986) Role of veins and cerebral venous pressure in disruption of the blood-brain barrier. Circ Res 59:216–20.

5. Chai PJ, Skaryla LA, Ungerleider RM, Greely VJ, Kern FH, et al. (1993) Jugular ligation does not increase intracranial pressure but does increase bitemporal cerebral blood flow and metabolism. Crit Care Med 23:1864–71.

6. Tovedal T, Jonsson O, Zemgulis V, Myrdal G, Thelin S, et al. (2010) Venous obstruction and cerebral perfusion during experimental cardiopulmonary bypass. Interact Cardiovasc Thorac Surg 11:561–6.

7. Levason JE, Wei EP, Raper AJ, Kontos AA, Patterson JL (1975) Detailed description of a cranial window technique for acute and chronic experiments. Stroke 6:308–17.

8. Plucinski J, Frydrychowski AF, Kaczmarek J, Juzwa W (2000) Theoretical foundations for non-invasive measurement of variations in the width of the subarachnoid space. J Biomed Opt 5:291–299.

9. Frydrychowski AF, Rojewski M, Guminski W, Kaczmarek J, Juzwa W (2001) Near infrared transillumination-back scattering (NIR-T/BSS) – a new method for non-invasive monitoring of changes in width of subarachnoid space and magnitude of cerebrovascular pulsation. Opto-Electron Rev 9:397–402.

10. Frydrychowski AF, Rojewski M, Guminski W, Kaczmarek J, Juzwa W (2002) Technical foundation for non-invasive assessment of changes in the width of the subarachnoid space with near-infrared transillumination-backscattering sounding (NIR-T/BSS). IEEE Trans Biomed Eng 49:887–904.

11. Plucinski J, Frydrychowski AF (2007) New aspects in assessment of changes in width of subarachnoid space with near-infrared transillumination/backscattering sounding, part 1: Monte Carlo numerical modeling. J Biomed Opt 12:044015.

12. Frydrychowski AF, Plucinski J (2007) New aspects in assessment of changes in width of subarachnoid space with near-infrared transillumination-backscattering sounding, part 2: clinical verification in the patient. J Biomed Opt 12:044016.

13. Li Z, Wang Y, Li Y, Wang Y, Li J, et al. (2010) Wavelet analysis of cerebral oxygenation signal measured by near infrared spectroscopy in subjects with cerebral infarction. Microvasc Res 80:142–7.

14. Li Z, Zhang M, Wang Y, Wang Y, Xin Q, et al. (2010) Wavelet analysis of sacral tissue oxygenation oscillations by near-infrared spectroscopy in persons with spinal cord injury. Microvasc Res 81:81–7.

15. Frydrychowski AF, Pankiewicz P, Sowiński P, Krzyżowski J (2009) Cerebrovascular pulsation and width of subarachnoid space during electroconvulsive therapy. J ECT 25:299–305.

16. Frydrychowski AF, Wszedybyl-Winklewska M, Bandurski T, Winklewski PJ (2011). Flow-induced changes in pial artery compliance registered with a non-invasive method in rabbits. Microvasc Res 82:156–162.

17. Frydrychowski AF, Wszedybyl-Winklewska M, Guminski W, Lass P, Bandurski T, et al. (2011) Effects of acute hypercapnia on the amplitude of cerebrovascular pulsation in humans registered with a non-invasive method. Microvasc Res 83:229–236.

18. Wszedybyl-Winklewska M, Frydrychowski AF, Winklewski P (2012) Assessing changes in pial artery resistance and subarachnoid space width using a non-invasive method in healthy humans during the handgrip test. Acta Neurobiol Exp 72:80–88.

19. Frydrychowski AF, Szarmach A, Czaplewski B, Winklewski PJ (2012) Subarachnoid space: new ways by an old dog. PLOS One: 7:e37529.

20. Frydrychowski AF, Wszedybyl-Winklewska M, Guminski W, Przyborska A, Kaczmarek J, et al. (2011) Use of Near Infrared Transillumination/Back Scattering Sounding (NIR-T/BSS) to assess effects of elevated intracranial pressure on width of subarachnoid space and cerebrovascular pulsation in animals. Acta Neurobiol Exp 71:313–321.

21. Winklewski PJ, Frydrychowski AF (2012) Cerebral blood flow, sympathetic nerve activity and stroke risk in obstructive sleep apnea. Is there a direct link? Blood Press DOI: 10.3109/000330510.2012.701467.

22. Bateman GA (2004) Pulse wave encephalopathy: a spectrum hypothesis incorporating Alzheimer’s disease, vascular dementia and normal pressure hydrocephalus. Med Hypotheses 62:182–7.

23. Bateman GA, Levi CR, Scholfield P, Wang Y, Lovent EC (2008) The venous manifestations of pulse wave encephalopathy: windkessel dysfunction in normal aging and senile dementia. Neuroradiology 50:491–7.

24. Chung CP, Hu HH (2010) Pathogenesis of leukoaraiosis: role of jugular venous reflex. Med Hypotheses 75:85–90.

25. Heinrich SS, Edwards J (1971) Vascular responses to acute intracranial hypertension. J Neurol Neurosurg Psychiatry 34:587–601.

26. Greitz D, Franck A, Norrild B (1993) On the pulsatile nature of intracranial and spinal CSF-circulation demonstrated by MR imaging. Acta Radiol 34:321–8.

27. Bateman GA (2000) Arterial inflow and venous outflow in idopathic intracranial hypertension associated with venous outflow stenoses. J Clin Neurosci 10:482–8.

28. Baumbach GL (1996) Effects of increased pulse pressure on cerebral arterioles. Hyperension 27:159–67.

29. Henkens JL, Kroon AA, van Oostenbrugge RJ, Gronenschild EH, Fuss-Lejune MM, et al. (2008) Increased aortic pulse wave velocity is associated with silent cerebral small-vessel disease in hypertensive patients. Hypertension 52:1120–6.

30. Hirata K, Yagimuna T, O’Rourke MF, Kawakami M (2006) Age-related changes in carotid artery flow and pressure pulses: possible implications for cerebral microvascular disease. Stroke 37:2552–6.

31. Zamboni P, Menegatti E, Weinstock-Gutmann B, Dwyer MG, Schirda CV, et al. (2011) Hyperperfusion of brain parenchyma is associated with the severity of chronic cerebrospinal venous insufficiency in patients with multiple sclerosis: a cross-sectional preliminary report. BMC Med 9:22.

32. Bateman GA (2000) Vascular compliance in normal pressure hydrocephalus. AJNR Am J Neuroradiol 21:1574–85.

33. Gisolf J, van Lieshout JJ, van Heusden K, Pott F, Stok WJ, et al. (2004) Human cerebral venous outflow pathway depends on posture and central venous pressure. J Physiol 560:317–27.

34. Zamboni P, Consotti G, Gallotto R, Gianselini S, MENEGATTI E, et al. (2009) Venous collateral circulation of the extracranial cerebrospinal outflow routes. Curr Neurol Neurosci Rep 9:204–12.

35. Wszedybyl-Winklewska M, Frydrychowski AF, Michalska BM, Winklewski PJ (2011) Effects of the Valsalva maneuver on pial artery pulsation and subarachnoid space in healthy adults. Microvasc Res 82:369–373.

36. Waigere BP, Gertsch S, Ammann RA, Pfenninger J (2003) Reproducibility of the blood flow index as noninvasive, bedside estimation of cerebral blood flow. Intensive Care Med 29:196–200.

**Author Contributions**

Conceived and designed the experiments: AFF PJW. Performed the experiments: AFF. Analyzed the data: PJW WG. Wrote the paper: AFF PJW WG. Statistical analysis: PJW.