The effects of sensory stimulation on postural stability after whiplash injury

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Ethical review	Approved WMO
Status	Pending
Health condition type	Other condition
Study type	Interventional

Summary

ID

NL-OMON32129

Source ToetsingOnline

Brief title Postural control after a whiplash accident

Condition

- Other condition
- Muscle disorders
- Spinal cord and nerve root disorders

Synonym neck sprain; neck strain

Health condition

symptomen als gevolg van whiplash trauma

Research involving

Human

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Sponsors and support

Primary sponsor: Vrije Universiteit Source(s) of monetary or material Support: Ministerie van OC&W

Intervention

Keyword: chronische pijn, postural control, sensory inormation, sensory integration therapy, whiplash

Outcome measures

Primary outcome

From the COP time-series, recorded by the force plate, we will calculate several standard parameters that are often used to quantify postural regulation, using in-house software (MATLAB). We will derive measures related to: 1) the overall amount of sway, 2) the variability and consistency of the sway, 3) the frequency contents of the signal, and 4) the smoothness of the signal.

For the three questionnaires we will calculate the total sum score of each measure.

All statistical analyses will be performed using SPSS version 14.0. A mixed factors ANOVA will be used. The between-subject factor is group and the within-subject factor is condition. We will conduct a separate ANOVA for the static balance conditions (A1 to A4) and the dynamic balance conditions (B1 to B4). Level of significance will be set to p <0.05.

Secondary outcome

Spearman correlations will be calculated between balance scores and

Study description

Background summary

Individuals with chronic neck and shoulder problems caused by whiplash injury not only suffer from pain in these areas, but they also often tend to have dizziness complaints, concentration problems, hypersensitivity to sensory information/stimulation, and a disturbed balance system. These complaints are collectively known in the literature as whiplash associated disorders (WAD). A crucial factor in disturbed balance appears to be that the different sensory modalities that are used for the regulation of balance are not properly tuned. A key aspect of rehabilitation is aimed at reducing the whiplash associated sensory complaints. This is known in therapeutic circles as sensory integration therapy (SI). However, at the moment little scientific knowledge exists on how the senses work together in the regulation of balance. The aim of this research is to investigate -using posturography- what the effects are of sensory manipulations on the regulation of balance in WAD patients. A group of 15 WAD patients and 15 unaffected controls will perform a number of simple balance tests. At the same time, movements of the body center of pressure will be recorded using a force plate.

Study objective

The aim of this research is to examine the effects of changes in proprioceptive and visual information on balance control in WAD sufferers. Although clinical experience suggests more effective balance control due to elevated proprioceptive information, this has never been quantified using objective posturographic measures. Balance will be measured by recordings of body sway, or more technically, the centre of pressure (COP) time-series registered with a force plate.

The following questions will be addressed:

* Is balance control in the WAD group less efficient than in a control group?
* In the WAD group, will balance control benefit from added proprioceptive input?

* In the WAD group, will balance control suffer from added visual input?

* What are the combined effects of the WAD syndrome and the availability of sensory signals on postural control?

Study design

Before the experiment begin participants sign the informed consent. Prior to the recording of the balance, participants (both patients and controls) are

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asked to fill out the Dutch equivalent of the following three questionnaires.
1) The Dutch version of the dizziness Handicap Inventory (DHI), which measures the subjective impact of dizziness on the everyday life of the participants.
2) The Tampa Scale for Kinesiophobia to assess pain related fear of movement. For unaffected controls the modified version of the TSK will be used.
3) The Beck Depression Inventory.

These questionnaires are standard use in clinical routine and scientific investigations of chronic pain patients, including WAD.

Upon arrival in the measurement lab and after the forms have been collected by the experimenter, patients are asked to indicate their level of actual experienced pain on a Visual Analogue Scale (VAS), on a 100-points scale (0 = no pain; 100 severe pain).

For the collection of the balance data, participants (patients and controls) will stand on a forceplate, adopting a natural and comfortable stance position, with the arms hanging relaxed alongside the body. Participants will be standing bare footed or wearing socks but no shoes or sandals, in order to maximize the quality of the posturographic measurements. In some conditions a belt weighing 500 g will be worn around the waist. This belt is part of the occupational therapist's toolbox when conducting sensory integration therapy.

Two different tasks will be performed. First, static balance capability (conditions A1 to A4) will be measured. During the static balance conditions participants will be asked to maintain quiet stance and to look straight ahead. In the experimental conditions the availability of visual and proprioceptive information is manipulated, as follows:

- A1: quiet standing (baseline)

- A2: quiet standing + 500 g waist belt

These 2 conditions are repeated, but participants will see a coloured spot (dot of approximately 5 mm) on a monitor in front of them that gives participants instantaneous visual information of the position (and changes therein) of their centre of pressure (COP). Thus, vision is added as an extra source of feedback for balance control. The instruction will be to keep this dot within the confines of a circle of a larger radius. This will result in 2 additional conditions:

- A3: quiet standing + visual feedback

- A4: quiet standing + 500 g waist belt + visual feedback

Second, dynamic balance (conditions B1 to B4) will be measured. During the dynamic balance conditions participants will be asked to periodically shift their weight from their left leg to their right leg and back again, which thereby results in medio-lateral displacements of their COP. In all conditions participants will receive visual feedback of their COP, similar to conditions A3/A4. In conditions B1 and B2 the task is to perform a set of rhythmic weight shifts so that the feedback dot on the screen oscillates rhythmically from left to right, thereby keeping the dot within the area enclosed by two vertical bars. - B1: weight shifting (baseline)

- B2: weight shifting + 500 g waist belt

Also, participants have to perform a task, in which they are asked to

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move the feedback dot between moving objects (slowly moving squares) and try not to *collide' with any of the blocks:

- B3: virtual collision avoidance

- B4: virtual collision avoidance + 500 g waist belt

The static balance (A1 to A4) will have a duration of 1 minute each. The dynamic balance conditions (B1 to B4) will have duration of 2 minutes each. Each condition will be repeated 2 times. Between each condition, participants can take a rest upon request, during which they are allowed to leave the force plate and sit down if they want. All participants will wear headphones to filter out possible background noise, as this might distract attention from the task. No verbal cues will be given to the participants during the tests. The static balance and dynamic balance conditions will be presented in a block, but within a block the conditions 1 to 4 will be presented in a random order.

Intervention

See Study design

Study burden and risks

During the experiment participants will have the opportunity to grab a handrail, located on the left and right side of the force plate, if they experience dizziness or discomfort. Subjects will also be closely monitored by an experimenter. Participants are free to take a small rest (or even abort the experiment) when they experience too much discomfort, fatigue, or dizziness. We expect hardly any instances of discomfort, fatigue, or dizziness due to the very low level of physical activity of the conditions and due to the low complexity of the visual signal.

Contacts

Public Vrije Universiteit

van der Boechorststraat 9 1081 BT Amsterdam Nederland **Scientific** Vrije Universiteit

van der Boechorststraat 9 1081 BT Amsterdam Nederland

Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age Adults (18-64 years) Elderly (65 years and older)

Inclusion criteria

Age 18+ -Patients are classified as having WAD grade II or III, according to the Quebec Task Force classification Eigibility to participate in the sensory integration treatment program of the RCA

Exclusion criteria

Other orthopaedic and/or neurolgical disorders, that are unrelated to WAD that might influence postural control.

Cognitive or mental impairments that might undermine the understanding of the task instructions.

Study design

Design

Study type:	Interventional
Intervention model:	Other
Allocation:	Non-randomized controlled trial
Masking:	Open (masking not used)
Control:	Active
Primary purpose:	Diagnostic

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Recruitment

NL	
Recruitment status:	Pending
Start date (anticipated):	01-08-2008
Enrollment:	30
Туре:	Anticipated

Ethics review

Approved WMO	
Date:	27-08-2008
Application type:	First submission
Review commission:	METC Amsterdam UMC

Study registrations

Followed up by the following (possibly more current) registration

No registrations found.

Other (possibly less up-to-date) registrations in this register

No registrations found.

In other registers

Register CCMO ID NL23767.029.08