

Application of the ELFI-TECH monitor for measurement of nociception

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Ethische beoordeling	Positief advies
Status	Werving nog niet gestart
Type aandoening	-
Onderzoekstype	Interventie onderzoek

Samenvatting

ID

NL-OMON23564

Bron

NTR

Verkorte titel

Elfitor 2

Aandoening

Nociception
Healthy volunteers
Hemodynamics

Nociceptie
Gezonde vrijwilligers
Hemodynamiek

Ondersteuning

Primaire sponsor: LUMC

Overige ondersteuning: Elfi-Tech Ltd.

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Validation ELFI-TECH monitor

Toelichting onderzoek

Achtergrond van het onderzoek

Background of the study

Accurate measurement of nociception during anesthesia remains a challenging task. Nociception, which is defined as the neural process of encoding and processing noxious stimuli (noxious stimuli are actually or potentially tissue damaging events), will elicit behavioral, autonomic and hormonal responses in conscious and unconscious individuals. Detection of behavioral responses during anesthesia is often difficult due to the use of sedatives and muscle relaxants. Hence, anesthesiologists rely on autonomic nervous system responses to assess the nociceptive level of the patient. Most anesthesia health care providers, if not all, use changes in heart rate and blood pressure as markers of the occurrence of acute nociceptive events. While these variables may suffice when intense nociceptive stimuli occur, mild and moderate stimuli are often not detected or detected too late.

In recent years various indices of nociception have been developed with varying success in actually detecting nociceptive events. These indices derive a numerical value from single variables (such as heart rate variability, skin conductance, skin vasomotor reflex, the electroencephalogram, pupil diameter) or a combination of signals. There are various new developments, one of which is the Nociception Level or NoL has recently been studied by us at LUMC (P13.069). We successfully studied the relationship between nociceptive stimuli and NoL during anesthesia and surgery. A completely new technology is the Elfitor device (Elfi-Tech, Rehovot, Israel), which is based on dynamic light scattering (DLS) and which continuously and non-invasively measures skin blood flow, skin blood flow variability, heart rate, heart rate variability, respiration, oxygen saturation, hemoglobin concentration, blood pressure and cardiac output. Additionally the system allows rheologic measurements from endothelial red blood cell interaction.

Objective of the study

In the current study we will apply the Elfitor and NoL and collect its derived parameters

during nociceptive stimulation in healthy volunteers without and with the administration of a low-dose opioid analgesic. Our study aim is to assess whether the various parameters derived from the Elfitor device correlate with nociceptive stimulation and standard hemodynamic parameters.

Study design

The subjects will arrive at 8 AM in the laboratory K5-120 and will receive an intravenous access line. Next they will be familiarized with the setup, pain tests, pain scoring and monitoring devices.

After a 30 min rest period the pain tests will be applied. Heat and electrical pain tests are performed in random order. The duration of the test sequence is about 2 hours. Then after another 3 min rest, the opioid remifentanyl will be infused using a target controlled infusion system (Remifusor, Glasgow University). The target blood concentration is 2 ng/mL. The complete sequence of testing will be repeated, again with the electrical and heat pain tests randomized in sequence.

Total duration of the study is 5 hours. Subjects will remain in the laboratory until the effects of remifentanyl are completely worn off (this usually takes about 20 min). Only fit subjects are allowed to leave the laboratory.

The following measurements will be obtained during pain stimulation:

1. Elfitor (positioned on the inside of second digit and outside of the wrist) ipsilateral from where pain is applied, preferably the non-dominant hand (this leaves the dominant hand for pain scoring). The data will be logged on a laptop PC and will be analyzed off-line. Most important parameters will be: heart rate, heart rate variability, blood flow, blood flow variability, plethysmogram, blood pressure, stroke volume, stroke volume variability, Mayer waves, and endothelial-red blood cell interaction.
2. Continuous non-invasive cardiac output and blood pressure using the Nexfin/finger cuff device, positioned on the fourth digit (ring finger) of the hand at which the Elfitor is placed. The Nexfin estimates finger systolic and diastolic blood pressure and cardiac output.

3. Pulse oximetry and ECG monitoring will be performed as safety measures.

4. NoL fingerprobe, positioned on the third digit of the non-dominant hand. The NoL combines information from several physiological parameters which represent different autonomic pathways. This information includes the photoplethysmogram, skin conductance level, fluctuations in skin conductance, and their time derivatives.

All participants will be familiarized with the study design, pain tests and scoring system. Pain intensity is scored using an 11-point numerical rating score (NRS) ranging from 0 (no pain) to 10 (maximum tolerated pain).

Test 1. Electrical pain is induced by placing electrodes on the tibial surface of the right leg. Electrical currents are applied using a locally designed and constructed computer interfaced current stimulator (CICS, Leiden University Medical Center, Leiden, The Netherlands). A preset increasing current at 0.5 mA/s is delivered and subjects are instructed to score the first time they feel pain (pain threshold, PTh) and the highest pain sensation they feel during the stimulation (pain tolerance, PTol) by pushing a button. This test is performed three times, 5-10 min apart.

Test 2. The values of PTh and PTol are used to construct a linear distribution of 8 interpolated currents. For example if PTh is 11 mA and PTol 20 mA, the interpolated temperatures are 12, 13, 14, 15, 16, 17, 18 and 19 mA. These 8 currents are subsequently presented in random order to the participants at 3-5 min intervals. All subjects are blinded to the sequence and intensity of the stimuli. They will be asked to rate the different stimuli on the 11-point NRS.

Heat pain stimulation.

An electronic Visual Analogue Scale (eVAS) will be used to quantify pain intensity in response to a noxious thermal stimulus. The thermal stimulus will be applied on the volar side of the forearm using a thermal probe (a 3 „e 3 cm thermode) of the TSA-II NeuroSensory Analyzer (Medoc Ltd, Ramat Yishai, Israel). This is a computer-controlled device capable of generating highly reproducible thermal stimuli. The eVAS will be measured electronically using a slide potentiometer (length = 10 cm) that can be moved from the left (0 or no pain) to the right (10 or most intense pain imaginable). Using a hand the subject can move the slide during the heat stimulator test. The eVAS is recorded and collected on disk for further analyses. In this study a constant increase in temperature will be applied, from 32 to 51 oC in 3 min after which the stimulus will be terminated, or at any time earlier when the eVAS score equals 10

cm. The test will be performed three times, 5-10 min apart; each time the thermode will be repositioned to prevent sensitization or adaptation.

20 healthy volunteers (10 males, 10 females) will be recruited to participate in the study. In case of early termination from the study, the subject will be replaced with a volunteer of the same sex.

Doel van het onderzoek

In the current study we will apply the Elfitor monitor and collect its derived parameters during nociceptive stimulation in healthy volunteers without and with the administration of a low-dose opioid analgesic. Our study aim is to assess whether the various parameters derived from the device correlate with nociceptive stimulation and standard hemodynamic parameters

Onderzoeksopzet

Testing will take place on one day and takes 5 hours

Onderzoeksproduct en/of interventie

Remifentanil infusion (blood target concentration 2ng/ml)
Pain testing

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

Subjects of either sex, aged 18-34 years with a body mass index < 30 kg/m²

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

Severe medical disease including hypertension, liver/renal disease, neurological disorders, diaphragmatic hernia/pyrosis;
(history of) psychiatric or neurological disease;
pregnancy/lactation; allergy to study medication;
(history of) illicit drug abuse/alcoholism

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Anders
Toewijzing:	N.v.t. / één studie arm
Blindering:	Enkelblind
Controle:	N.v.t. / onbekend

Deelname

Nederland	
Status:	Werving nog niet gestart
(Verwachte) startdatum:	01-09-2015

Aantal proefpersonen: 20
Type: Verwachte startdatum

Ethische beoordeling

Positief advies
Datum: 07-09-2015
Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL5230
NTR-old	NTR5454
Ander register	METC LUMC : P15.156

Resultaten