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Small and large cutaneous fibers display different excitability properties to slowly increasing ramp pulses

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Title: Small and large cutaneous fibers display different excitability 1 properties to slowly increasing ramp pulses 2 3 assessed with the perception threshold tracking technique 4 Authors: Jenny Tigerholm², Tatiana Nielson Hoberg, Dorthe Brønnum^{1,3}, Mette 5 Vittinghus^{1,4}, Ken Steffen Frahm^{1,2}, Carsten Dahl Mørch^{1,2*}. 6 7 8 (1) SMI, Department of Health Science and Technology, Aalborg University, Fredrik Bajers 9 Vej 7 D3, 9220 Aalborg, Denmark. 10 (2) Integrative Neurodcience group, CNAP - Center for Neuroplasticity and Pain, SMI, 11 Department of Health Science and Technology, Aalborg University, Fredrik Bajers Vej 7 D3, 12 9220 Aalborg, Denmark. 13 (3) Centre for Clinical Research, North Denmark Regional Hospital, Bispensgade 37, 9800, 14 Hjørring, Denmark 15 (4) It-center for Telemedicin, Region Midtjylland, Oluf Palmes Allé 36, 8200 Aarhus N, 16 Denmark 17 18 19 Acknowledgements 20 Funded by Center for Neuroplasticity and Pain (CNAP). CNAP is supported by the Danish 21 National Research Foundation (DNRF121). 22 *Corresponding Author: 23 Carsten Dahl Mørch, 24 Integrative Neuroscience group, CNAP

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Running title: Accommodation in cutaneous nerves

Abstract

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33 The excitability of large nerve fibers is reduced when their membrane potential is slowly 34 depolarizing, i.e. the fibers display accommodation. The aim of this study was to assess 35 accommodation in small (mainly $A\delta$) and large ($A\beta$) cutaneous sensory nerve fibers using the 36 perception threshold tracking (PTT) technique. 37 38 Linearly increasing ramp currents (1 ms -200 ms) were used to assess the excitability of the 39 nerve fibers by cutaneous electrical stimulation. To investigate the PPT technique's ability to 40 preferentially activate different fiber types, topical application of lidocaine/prilocaine 41 (EMLA) or a placebo cream was applied. By means of computational modelling, the 42 underlying mechanisms governing the perception threshold in the two fiber types was studied. The axon models included the voltage-gated ion channels: Na_{TTXs} , Na_{TTXr} , Na_P , K_{Dr} , 43 44 K_M , and HCN. 45 46 Large fibers displayed accommodation, whereas small fibers did not display accommodation 47 (p<0.05). For the pin electrode, a significant interaction was observed between cream (EMLA 48 or placebo) and pulse duration (p<0.05) whereas for the patch electrode, there was no 49 significant interaction between cream and duration which supports the pin electrode's 50 preferential activation of small fibers. The results from the computational model suggested 51 that 52 differences in accommodation between the two fiber types may originate from selective 53 expression of voltage-gated ion channels, particularly the transient Na_{TTXr} and/or K_{Dr}. 54

- 55 The PTT technique could assess the excitability changes during accommodation in different
- 56 nerve fibers. Therefore, the PTT technique may be a useful tool for studying excitability in
- 57 nerve fibers both in healthy as well as in pathological conditions.

- 59 Keywords: Accommodation, perception threshold tracking technique, nerve fiber excitability,
- 60 voltage-gated ion channels, multi-compartmental model

New & Noteworthy

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When large nerve fibers are stimulated by long, slowly increasing electrical pulses interactive mechanisms counteract the stimulation, which is called accommodation. The perception threshold tracking technique was able to assess accommodation in both small and large fibers. The novelty of this study is that large fibers displayed accommodation, whereas small fibers did not. Additionally, the difference in accommodation between the fiber could be linked to expression of voltage-gated ion channels by means of computational modeling.

Introduction

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| Neuropathic pain has a prevalence rate of 6.9 % - 10 % (van Hecke, et al., 2014) and |
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| manifests itself through symptoms of burning pain, shooting pain, allodynia, and |
| hyperesthesia (Hovaguimian & Gibbons, 2011). The underlying mechanisms of small fiber |
| neuropathy are unknown, but altered excitability has been detected both in patients with |
| peripheral neuropathic pain as well as in animal models of neuropathy (Serra, et al., 2012; |
| Serra, et al., 2011). Tactile information is passed through afferents with larger diameters (Aß |
| fibers), whereas nociceptive information is passed through small diameter afferent fibers (C |
| and $A\delta$ fibers). The main obstacle when studying small fibers in humans is their high |
| electrical activation thresholds which makes it technically challenging to study the fibers in |
| isolation without activation of the large fibers. To overcome this obstacle, electrodes with |
| small cathodes (pin electrodes) have been used (Bromm & Meier, 1984; Nilsson & |
| Schouenborg, 1999; Kaube, et al., 2000; Inui, et al., 2002; Klein, et al., 2004; Otsuru, et al., |
| 2009; Lelic, et al., 2012; Hennings, et al., 2017). These Pin electrodes preferentially activate |
| the superficial small fibers by generating a high current density in epidermis and thereby |
| avoiding activating the large fibers which are terminating in dermis (Hilliges, et al., 1995; |
| Ebenezer, et al., 2007; Provitera, et al., 2007; Mørch C, et al., 2011; Myers, et al., 2013). |
| Estimations of nerve fiber conductance velocity support the non-invasive pin electrode's |
| preferential activation of small fibers (Inui, et al., 2002; Otsuru, et al., 2009; Lelic, et al., |
| 2012). Test subjects perceive the pin electrode stimulation as needle pricking, stabbing and |
| sharp, and distinctly different from large cathode electrodes (patch electrodes) designed to |
| activate large fibers (Bromm & Meier, 1984; Hugosdottir, et al., 2017; Lelic, et al., 2012). A |
| low stimulation intensities, non-invasive pin electrodes preferentially activate the thinly |

95 myelinated small A\delta fibers (Inui, et al., 2002; Lelic, et al., 2012), whereas the invasive pin 96 electrode may also activate the unmyelinated C fibers (Otsuru, et al., 2009). 97 98 Our research group has developed the Perception Threshold Tracking (PTT) technique, 99 which assesses neuronal excitability by measuring the perception threshold to cutaneous 100 electrical stimulation, using different pulse shapes and durations (Hennings, et al., 2017; 101 Hugosdottir, et al., 2017). The perception threshold is defined, in the current study, as the 102 intensity of the stimulus required for the subject to perceive the stimulus at the site of 103 stimulation. In a recent study, the strength-duration properties and threshold electrotonus 104 were assessed for both small and large fibers (Hennings, et al., 2017). Interestingly, the 105 excitability assessments differed between small and large fibers indicating different 106 membrane properties between the two fiber types. In this study, the objective was to study the 107 excitability to linearly increasing cutaneous ramp stimulation in small and large fibers. For 108 motor fibers, the excitability of both small and large diameter motor neurons has been shown 109 to be reduced for long ramp pre-pulses (Hennings, et al., 2005). When large nerve fibers are 110 stimulated by long, slowly increasing electrical stimulation (50-200 ms) a higher stimulation 111 current is needed to activate the fiber compared to a shorter electrical stimulation (Lucas, 112 1907; Kugelberg, 1944) (see figure 1A). This altered excitability will in the current study be 113 referred to as fiber accommodation (Kugelberg, 1944). The ability of accommodation of 114 each nerve fiber is determined by the intrinsic properties of its membrane (Stoney S & 115 Machne, 1969). Particularly, the resting membrane potential has been shown to modulate the

accommodation in large fibers (Baker & Bostock, 1989; Bostock, et al., 1991). The strong

influence of the resting membrane potential on accommodation indicates that voltage-gated

ion channels play a significant role in generating accommodation. From patch clamp and

threshold electrotonus experiments in rats, potassium channels have been identified to alter

the accommodation by rectification of the membrane potential (Baker & Bostock, 1989;

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| Stoney S & Machne, 1969). However, a computational study has shown that almost any |
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| alteration of the density of the voltage-gated ion channel will alter the accommodation, |
| whereby the parameter which had the strongest influence was the inactivation of transient |
| sodium channels (Frankenhaeuser & Vallbo, 1965). Since small and large fibers have |
| different expressions of voltage-gated ion channels (Akopian, et al., 1996; Gold, et al., 1996 |
| Djouhri, et al., 2003; Gao, et al., 2012), the hypothesis is that their respective |
| accommodations may differ. For instance, small fibers express two TTX resistant sodium |
| channels (Na _v 1.8 and Na _v 1.9), which are lacking in large fibers (Akopian, et al., 1996; |
| Djouhri, et al., 2003). Therefore, the purpose of this study was to measure the |
| accommodation in small fibers as well as large fibers, and link the accommodation to |
| membrane properties. |
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Materials and methods

This is a combined clinical and computational study (see figure 1). In the experimental study, the accommodation in small and large cutaneous nerve fibers was assessed using the PTT technique. For long ramp pulses (>20 ms) the excitability can be reduced, and the intensity needed for the subject to perceive the stimulus is increased, this is denoted as accommodation in the current study. Topical application of EMLA cream was used to validate the pin electrode's preferential activation of small fibers. To identify the possible mechanisms for the different accommodation between the two fiber classes, two multi-compartment models were developed.

Experimental study

Subjects

20 healthy subjects participated in the study, however one subject was excluded due to technical issues, thus data analysis was completed for 19 subjects (10 males, 9 females, age 34.6±13.3 years). The subjects were given detailed written and verbal information and signed an informed consent form prior to participation. The study was approved by the local ethics committee (Den Videnskabsetiske Komité, Region Nordjylland, approval number: N-20120046) and conducted according to the declaration of Helsinki. Exclusion criteria were; a) addiction or prior addiction to cannabis, opioids, or other drugs, b) skin diseases, c) infectious diseases, d) conditions that might lead to peripheral neuropathy, and e) pain relieving medication within the last 48 hours.

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Experimental setup

The experiment consisted of one experimental session lasting 3.5 hours. The subjects were placed in a comfortable inclined position in a hospital bed throughout the session.

Two surface electrodes were used to preferentially activate small and large fibers as described in previous papers (Hennings, et al., 2017). A cutaneous pin electrode with a circular array of 16 small area cathodes made of blunted stainless steel with a diameter of 0.2 mm protruding 1 mm from the base of the electrode and a concentric stainless steel disc anode with an area of 8.8 mm was used to preferentially activate small fibers (see figure 1B). A large area surface AgAgCl cathode (Patch, 20 x 15 mm; Neuroline 700; Ambu A/S, Ballerup, Denmark) in combination with a larger anode (5 x 9 cm; Pals Neurostimulation Electrode Axelgaard, CO., Ltd., California) was used to preferentially activate large fibers.

PTT was performed with a computer-controlled program, LabBench (SMI®, Aalborg University, Denmark) stimulating with a DS5 Isolated Bipolar Current Stimulator (Digitimer Ltd, Letchworth Garden City, UK). Perception thresholds were assessed by an adaptive staircase method; Stimuli were given with an inter-stimulus interval of 1 second and increased by 15% of the last stimulus until the subject indicated perception by pressing a handheld response button (SMI®, Aalborg University, Denmark). After indicating perception by pushing the button, the stimulation intensity was maintained two consecutive times and if these stimulations were also perceived, the intensity was decreased by 15% until the stimulation was not perceived anymore, as indicated by not responding three consecutive times. In the four following sequences the intensity increased and decreased by 7.5%, 3.5%,

3%, and 3%. The perception threshold was calculated as a weighted average of all 10 measurements. Accommodation was assessed by linearly increasing ramp currents of 1 ms, 10 ms, 25 ms, 50 ms, 100 ms and 200 ms. The order of the electrical stimulations was randomized in a single-blinded manner.

To validate that the pin electrode preferentially activates the small fibers, topical application of EMLA was used to block the small fibers (which should cause increase activation threshold) (Bjerring & Arendt-Nielsen, 1990). A cream similar in color, smell, and consistency was used as placebo. The creams were supplied in identical vials and the order of the creams (EMLA or placebo) was randomized 1:1. The experimenter applied 4 grams of cream (either EMLA or placebo) to a 5 x 5 cm skin area on the volar part of the forearm 5 cm distal from elbow pit under an impermeable plastic occlusive film. After 60 minutes the cream was removed from the first arm and a cream was applied similarly to the opposite forearm. Testing began on each arm 30 minutes after removal of the cream to ensure a stable effect of EMLA during performance of the perception threshold measurements.

Data analysis

All statistical calculations were performed using MATLAB 2016b (MathWorks, Natick, Massachusetts, USA) and SPSS 25 (IBM SPSS Statistics, Armonk, New York, USA). To obtain normality, the perception thresholds were normalized to the threshold for the 1 ms duration pulse and log transformed. A three-way repeated measures ANOVA was carried out to analyze the difference between the electrodes (pin or patch), the cream (EMLA and placebo) and the ramp pulse durations (1 ms, 10 ms, 25 ms, 50 ms, 100 ms and 200 ms). In

the event of significant interactions involving the electrodes, a two-way repeated measures ANOVA was carried out for each electrode with cream (EMLA and placebo) and ramp pulse durations (1 ms, 10 ms, 25 ms, 50 ms, 100 ms and 200 ms). The Greenhouse-Geisser method was used to adjust for non-spherical covariance matrices. Pairwise comparisons of the estimated marginal means were corrected for multiple-comparison with the Sidak method when comparing differences between ramp pulse durations. Statistical significance was defined as p < 0.05.

Computational model

Two computational fiber models were developed in the simulator environment NEURON ((Hines & Carnevale, 1997), version 7.6); one myelinated fiber model ($A\beta$ model) representing a large fiber and one unmyelinated fiber model representing the unmyelinated intraepidermal part of an $A\delta$ fiber ($A\delta$ model). The cutaneous electrical stimulation will activate a population of fibers, but in order to reduce the computational complexity, one fiber will represent the mean of a population of fibers. No sensory transductions were modeled, because during electrical stimulation an action potential is generated by shifting the voltage across the cell membrane and no sensory transduction within the sensory terminal is occurring. Instead of a sensory terminal, the $A\beta$ model terminated in a node of Ranvier. All morphological parameters are listed in table 1. The number of compartments were 5000 for the $A\delta$ model and 5133 for the $A\beta$ model (node of Ranvier = 3 compartments, internode = 500 compartments and Juxtaparanode= 5 compartments) and the equations were solved using the variable time step method in NEURON. The resting membrane potential was set to -60 mV (Fang, et al., 2005). The delayed rectifier potassium channels' voltage dependency was shifted 15 mV towards hyperpolarization in order to generate a rectification of the action

potential. Four sodium channels were implemented: two transient TTX- sensitive sodium currents (Na_{TTXs}), the transient TTX resistant sodium current (Na_{TTXr}) and the persistent sodium current (Na_P).

The morphological parameters are listed in table 1.

Table 1. Parameters of the morphology for the fiber models

| | A8 model | Aβ model | Reference |
|----------------------------|------------|-----------------------|---------------------------|
| Diameter | 0.5-3.5 μm | 9 μm | |
| Nodal length | | 3 μm | (Berthold & Rydmark, |
| | | | 1983) |
| Internodal length | | 500 μm | (Nilsson & Berthold. , |
| | | | 1988; Provitera, et al., |
| | | | 2007) |
| Juxtaparanodal length | | 5 μm | (Poliak, et al., 2003) |
| Capacitance nodal/branch | | 1 μF/cm ² | (Amir & Devor, 2003) |
| Capacitance myelin | | $0.0141 \ \mu F/cm^2$ | (Amir & Devor, 2003) |
| | | | C=1/(myelin sheet +1) |
| Resting membrane potential | -60 mV | -60 mV | (Fang, et al., 2005) |
| Number of myelin sheets | | 70 | (Provitera, et al., 2007; |
| | | | Berthold & Rydmark, |
| | | | 1983) |
| Intra cellular resistance | 130 Ωcm | 130 Ωcm | |
| Total model length | 5000 μm | 5133 μm | |

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| 244 | Table 2. The models of voltage-gated ion channels. K_{Dr} : delayed rectifier potassium channel, |
| 245 | K_{M} : slow potassium channel, HCN: hyperpolarization-activated current, Na_{TTXs} : TTX |
| 246 | sensitive current, Na _{TTXr} : TTX resistant current, Na _P : persistent sodium current. |
| 247 | |
| 248 | |

| Aδ model | Model Reference | Spatial location | Maximal |
|--|-----------------------------|------------------|----------------------------------|
| | | | Conductance (S/cm ²) |
| Na _{TTXr} | (Tigerholm, et al., 2014) | Unmyelinated | 0.0435 |
| (Na _v 1.8) | | nerve | |
| Na _p | (Tigerholm, et al., 2014) | Unmyelinated | 3.5549x10 ⁻⁵ |
| (Na _v 1.9) | | nerve | |
| Na _{TTXs} | (Tigerholm, et al., 2014) | Unmyelinated | 0.0166 |
| (mainly Na _v 1.7) | | nerve | |
| K_{Dr} | (Tigerholm, et al., 2014) | Unmyelinated | 3.4023x10 ⁻⁴ |
| | | nerve | |
| K _M | (Tigerholm, et al., 2014) | Unmyelinated | 1.0460x10 ⁻⁶ |
| | | nerve | |
| HCN | (Tigerholm, et al., 2014) | Unmyelinated | 1.4275x10 ⁻⁶ |
| | | nerve | |
| Aβ model | | | |
| Na _{TTXs} (Na _v 1.6) | (Watanabe, et al., 2002) | Nodes of Ranvier | 0.4394 |
| Nap | (Jankelowitz, et al., 2007) | Nodes of Ranvier | 7.7731x10 ⁻⁵ |
| K _{Dr} | (Tigerholm, et al., 2014) | Juxtaparanode | 0.0065 |
| HCN | (Tigerholm, et al., 2014) | Juxtaparanode | 9.1358x10 ⁻⁴ |
| | | | |
| K _M | (Tigerholm, et al., 2014) | Nodes of Ranvier | 0.0021 |
| Leak channel | (Tigerholm, et al., 2014) | Internode | 1.0000x10 ⁻⁷ |
| | | | |

250 Table 3. Action potential characteristics

| Aδ model | Aβ model |
|----------|----------|
| | |

| Action potential height | Initiation: 123 ms | Initiation: 127 ms |
|-------------------------|--------------------------|--------------------------|
| | End of the model: 120 ms | End of the model:114 ms |
| Action potential width | Initiation: 3.7 ms | Initiation: 1.6 ms |
| | End of the model: 3.6 ms | End of the model: 1.5 ms |
| Velocity | 0.54m/s | 11m/s |

A8 model

The $A\delta$ model was developed by modifying a previously published, detailed multi-compartment model of a C-fiber (Tigerholm, et al., 2014; Tigerholm, et al., 2015). In this study, the C-fiber model has been simplified by the removal of the ion concentration dynamics. The diameter was increased for the C-fiber model to be consistent with an $A\delta$ -fiber's morphology. The $A\delta$ model consists of two sections. The first section (500 μ m) starts with a diameter of 0.5 μ m, which is increased linearly to 3.5 μ m in diameter. The second section is a cylinder with a diameter of 3.5 μ m connecting to the first section (see figure 1C). The $A\delta$ -model did not include any myelination since only the superficial part of the $A\delta$ fiber was modeled, and the $A\delta$ fibers lose their myelin when entering the epidermis (Provitera, et al., 2007). One TTX-sensitive sodium channel (Na_{TTXs}, mainly Na_v 1.7) and two resistant sodium channels were implemented (Na_v 1.8 and Na_v 1.9). Additionally, two potassium channels and one HCN channel were implemented. See table 2 for the voltage-gated ion channel model references for the equations of the steady-state parameters, and their time-constants. The equations describing the ion channel dynamics are stated in the supplemental data. The action potential characteristics are presented in table 3.

Aß model

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The Aβ model consists of three different morphological sections: nodes of Ranvier, Juxtaparanode and myelinated fiber (see figure 1C). The three different sections of the membrane have different electrical properties as well as distribution of voltage-gated ion channels (see table 1 and 2). The capacitance of the myelin section of the axon is dependent of the thickness of the myelin i.e. the number of myelin sheets (Amir & Devor, 2003). The capacitance in the current study was calculated by C=1/(the number of myelin sheets +1), which is a method adopted from the Amir and Devor study (2003). One TTX-sensitive sodium channel (Na_{TTXs}, Nav 1.6) and one persistent sodium channel (Na_P) were implemented. Additionally, the two potassium channels and the HCN channel which were implemented in the A δ model was also implemented in the large fiber model. See table 2 for the voltage-gated channel model references for the equations of the steady-state parameters and their time-constants. The equations describing the ion channel dynamics are stated in the supplemental data. https://doi.org/10.5281/zenodo.3975475. The action potential characteristics are presented in table 3. The internodal distance is set to 500 µm, which is one quarter of the internodal distance measured in deeper layers (Nilsson & Berthold., 1988) since the internodal length of the nerve fibers is reduced when they enter superficial layers of the skin (Provitera, et al., 2007).

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Extracellular stimulation

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To simulate the cutaneous stimulation, the extracellular potential at the most superficial section of the fiber models was changed with the same shape as the current pulse applied

| through the electrode in the experiment. For the $A\delta$ model, the extracellular potential was |
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| altered for a section of 500 μm through the built-in function of the extracellular in NEURON. |
| For the $A\beta$ fiber model, the extracellular potential of the first node of Ranvier was altered. |
| The shape of the extracellular alteration was a ramp pulse with the same durations as were |
| tested in the experimental study. The extracellular potential was increased until the fiber |
| model generated an action potential (membrane potential higher than 0 mV) which |
| propagates to the end of the fiber model. |
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Constraints of the computational models

- The maximum conductances of the voltage-gated ion channels were defined by the following constraints:
- 309 1) For increasing durations of the ramp simulation, in the interval 50 ms 200 ms, the
 310 activation threshold of the Aδ model should be declining or constant.
- 311 2) For increasing durations of the extracellular potential in the interval 50 ms 200 ms,
 312 the activation threshold should increase for the Aβ model.
 - 3) If an action potential is generated (membrane potential > 0 mV), it should propagate to the end of the nerve fiber model
- 315 4) The Na_{TTXr} current should generate the action potential in the A δ model (Blair & Bean, 2002)
- 5) The action potential should be higher than 0 mV at the last section of the model for the simulation to be classified as a successful propagation.
- 319 6) The HCN maximum conductance should be at least two times higher in the $A\beta$ model 320 than in the $A\delta$ model (Gao , et al., 2012)

Results

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Small and large fibers have different accommodation to ramp pulse electrical stimulation

The absolute value of the perception threshold was higher for the patch electrode than the pin electrode across all durations of the ramp stimulations (figure 2). This was a direct consequence of the different cathode configurations. More interestingly, a significant difference of log-transformed and normalized thresholds was observed between the electrodes (p < 0.05) and interactions between cream (EMLA and placebo) and electrodes (p < 0.01) as well as between electrode and pulse duration (p < 0.001) indicating that accommodation was different between the two electrodes (see figure 2C). The electrodes were therefore analyzed individually.

For the patch electrode, significantly different thresholds were observed between the pulse durations (p < 0.001), but not between cream (p = 0.06), and there was no interaction between cream and duration. Comparing the estimated marginal means of the pulse durations showed that the thresholds for the 1 ms, 100 ms and the 200 ms pulses were higher than the thresholds to 20 ms, 50 ms pulses (p < 0.05; figure 2C), indicating that large nerve fibers accommodated to long duration ramp pulses.

For stimulation with the pin electrode in the EMLA arm, the 1 ms pulse threshold was significantly higher than the threshold to the 50 ms pulse (p < 0.05). No other significant threshold differences were observed between the 1 ms pulse and any of the other pulse durations, indicating similar, though not as pronounced, accommodation as large nerve fibers

activated by the patch electrode. Thus, the nerve fibers activated under this condition are probable not only large fibers, but a combination of both small and large fibers.

For stimulation with the pin electrode in the placebo arm, the 1 ms pulse threshold were significantly higher than all the longer pulses (p<0.05), indicating the small fibers activated by the pin electrode did not accommodate to ramp pulses.

The accommodation generated in the computational model

A wide range of voltage-gated ion channels with different dynamic properties were implemented in the computational model. In figure 3, the ionic currents during an action potential is illustrated. In figure 4A, the extracellular potential alteration needed to generate an action potential for different duration of the ramp pulses is illustrated. For a 1 ms pulse duration, the increase of the extracellular potential needed to activate the $A\delta$ model was 4.3 times larger than the $A\beta$ model. This is consistent with the high electrical stimulation intensity needed to activate small fibers with the patch electrode. In figure 4B, the relative extracellular potential alteration needed to generate an action potential is illustrated. The excitability of the two nerve fiber models was affected differently by long ramp pulses (see figure 4 C-F). For the $A\delta$ model, the membrane became more excitable when the membrane potential was increasing slowly (see figure 4C and 4E). For the 200 ms slowly-increasing stimulation the small fiber model even starts to produce two spikes at the end of the stimulation. For the $A\beta$ model, slow depolarization leads to reduced excitability and the inability to generate an action potential when the membrane is too depolarized (see figure 4D and 4F).

The influence of voltage-gated ion channels on accommodation

The influence of subtypes of voltage-gated ion channels on the accommodation is illustrated in figure 5. For the A β fiber model, an alteration of the Na_{TTXs} channel had the strongest influence on the accommodation, which can be explained by the high channel density in the node of Ranvier. However, changing any of the ion channel densities influenced the accommodation curve in the A β model. For the A δ model, changing the maximum conductance of subtypes of ion channels did not alter the accommodation substantially. Furthermore, when both the potassium channels' conductances were reduced with 50%, the A β model did not display any accommodation (see figure 5B, green dotted line). The purpose of this simulation was to evaluate the consistency between the behavior of the computational model and the experimental results, showing no accommodation when the potassium channels were blocked in an animal study (Stoney S & Machne, 1969).

To further analyze the influence of voltage-gated ion channels, larger perturbation of the maximum conductance was implemented in the computational model (see figure 6). When the maximum delayed rectifier conductance was reduced by 60%, the $A\beta$ model did not display any accommodation (see figure 6A-B, blue line). If instead the maximum conductance of delayed rectifier potassium was increased by 300% in the $A\delta$ model, the model displayed accommodation in a similar fashion as the $A\beta$ model (see figure 6C-D, blue dashed line). If instead the slow dynamic voltage-gated ion channels (HCN, Na_P , and K_M) were removed from the nerve fiber models, the accommodation was marginally altered (see figure 6, green lines). Interestingly, if all voltage-gated ion channels except the Na_{TTXs}

channel were removed from the $A\beta$ model, the model could still generate accommodation (see figure 6, light blue lines).

The influence of inactivation of sodium on accommodation

The inactivation of the transient sodium channels for the two models are compared in figure 7 where the extracellular potential was altered up to 10 mV and with a duration of 200 ms ramp pulse. The Na_{TTXs} channels in the $A\beta$ model were 84 % inactivated while the Na_{TTXr} channels in the $A\delta$ model were only 65% inactivated (see figure 7B). Despite that 10 mV alteration of the extracellular potential depolarizes the $A\beta$ model less than the $A\delta$ model (see figure 7A).

To further study, the influence of inactivation of sodium on accommodation, the steady-state inactivation curves of either Na_{TTXs} ($A\beta$ model) or the Na_{TTXr} ($A\delta$ model) were shifted. To compensate for the general excitability, the maximum conductance of the Na_{TTXs} was increased ($A\beta$ model), or the Na_{TTXr} was adjusted ($A\delta$ model), whereby the activation threshold for the 1 ms ramp stimulation remained within 5 % compared to the control model (no shifts of the inactivation). By shifting the steady-state inactivation curves, the accommodation could be generated in the $A\delta$ model (dashed lines, Figure 7E-F) and removed in the $A\beta$ model (dashed lines, Figure 7C-D).

Discussion

In this study, accommodation in small and large sensory fibers was estimated by the PTT technique. The PPT experiment showed that large fibers displayed accommodation to long ramp stimulation pulses, while small fibers did not. Furthermore, the results from the computational model suggested that the selective expression of voltage-gated ion channels may account for the difference in accommodation between the two fiber types.

Accommodation to ramp electrical stimulations differed between small and large fibers

Accommodation of nerve fibers has mainly been studied in humans by assessing the compound action potential in large fibers (Baker & Bostock, 1989; Bostock, et al., 1998; Kiernan, et al., 2000). Assessment of the compound action potential can be performed on large fibers but this is technically challenging to detect in small fibers. Therefore, our research group has previously used the perception threshold instead of compounded action potential for studying the strength-duration relationship and the threshold electrotonus (Hennings, et al., 2017; Hugosdottir, et al., 2017). In this study, accommodation has been studied in small and large fibers with this PTT technique. Accommodation of the median

nerve has previously been estimated by assessing the motor compound action potential (Hennings, et al., 2005). The threshold of the median nerve for 200 ms was 88%-96% of the threshold for the 1 ms duration. In our study, the threshold of the large fibers for 200 ms was 61% of the threshold of 1 ms duration. The lower value measured in our study could be explained by the difference in fiber types, motor vs sensory fibers. This is supported by the classic work of Kugelberg showing that motor fibers have more pronounced accommodation properties than the sensory fibers (Kugelberg, 1944). Interestingly, small fibers did not display accommodation i.e. had no significant increase in perception threshold for long ramp simulation. In our previous study, threshold electrotonus was studied for both large and small sensory fibers, and the result showed no significant differences between the perception thresholds of the two fiber types except for long (80ms) hyperpolarizing prepulses (Hennings , et al., 2017). The threshold electrotonus protocol assesses the effect of altering the membrane potential of nerve fibers activation, which is another method for probing accommodation. The discrepancy between the current study and our previous study could be explained by the fact that only a small prepulse was used.

Moreover, the ability of accommodation in nerve fibers is essential as it attempts to counteract the effect of a sustained stimulus, thereby limiting the generation of action potentials and repetitive firing from the neurons (Baker & Bostock, 1989). In the computational model of small fibers, multiple spikes were generated when a 200 ms ramp stimulation was applied (see figure 4C). Human and animal models of neuropathic pain states showed spontaneous activity or duplets when small fibers were stimulated (Serra, et al., 2012; Serra, et al., 2011). A possible explanation for this could be that the increased excitability occurring in these pain states leads to a depolarized membrane and thereby an enhanced possibility of multiple spike generation in the small fibers.

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Different expression of voltage-gated ion channels may generate the difference in accommodation between the two fiber types.

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The underlying mechanisms of accommodation have been studied in both threshold tracking experiments, DRG patch clamp experiments as well as in computational models (Baker & Bostock, 1989; Stoney S & Machne, 1969; Frankenhaeuser & Vallbo, 1965; Hennings, et al., 2005). It is clear that voltage-gated ion channels play an important role in generating the accommodation since altering the membrane potential or blocking potassium channels has a strong influence on accommodation (Bostock, et al., 1991; Baker & Bostock, 1989; Baker M, et al., 1987). However, which subtypes of the voltage-gated ion channels have the most substantial influence on the accommodation is still unclear. The ion channel parameter which had the most substantial influence on accommodation in a computational study was the inactivation of the sodium channel (Frankenhaeuser & Vallbo, 1965). However, almost all ion channel parameters altered in the computational model had a strong influence on the accommodation (Frankenhaeuser & Vallbo, 1965). This may explain why blocked potassium channels lead to almost a complete loss of accommodation in a threshold electrotonus experiment in rats (Baker & Bostock, 1989). This behavior could be reproduced in our AB fiber model when the maximum conductance of the potassium channels was reduced by 50% (see figure 5D). One of the most interesting studies of accommodation is a patch clamp study in which the excitability to an intracellular current stimulation in large DRG somas is studied (Stoney S & Machne, 1969). In that study, the membrane potential plateaued for a long ramp current stimulation and the action potential was not generated at the end of the stimulation but instead in the middle of the stimulation. This is consistent with the behavior displayed by the computational model of the $A\beta$ fiber where the action potential was generated at 80 ms for a ramp pulse of 200 ms (see figure 4D and 4F).

Furthermore, a support for the sodium inactivation important contribution to accommodation is the result that accommodation could be generated in the computational model when only TTX-sensitive sodium channel was implemented in the A β model (see figure 6B). These results suggest that the difference in inactivation between the Na_{TTXr} and Na_{TTXs} channels may contribute to generate the difference in accommodation measured between small and large fibers. The Na_{TTXr} sodium channel (Na_v 1.8) is only expressed in small fibers (Djouhri, et al., 2003) and the channel is inactivated at more depolarized membrane potential (Inactivation V_{1/2} =-32 mV (Blair & Bean, 2002)) than the Na_{TTXs} sodium channel (Na_v1.6), which is expressed in large fibers (Inactivation V_{1/2} =-55 mV (Smith, et al., 1998; Caldwell, et al., 2000)).

Clinical implications for perception threshold tracking

Available diagnostic tools to determine small fiber functionality mainly include clinical examination, skin biopsies and quantitative sensory testing. Such available diagnostic tools for small fiber neuropathy are insufficient to explore the mechanisms underlying neuropathy and there is a need for new methods (Smith, et al., 2017). Compared to the existing method for probing excitability in small fibers (microneurography) the PTT technique is an inexpensive and non-invasive method which makes it suitable for clinical settings. Whereas the PPT technique is a newly developed method and the diagnostic accuracy of PPT

techniques has not been evaluated, this study, as well as our previous studies (Hennings, et al., 2017; Hugosdottir, et al., 2017), have shown that the PTT technique may potentially be able to distinguish between membrane properties of both small and large nerve fibers. Further development of both the PPT method and the fundamental understanding of membrane currents related to neuropathy is required, but the PTT technique certainly has the possibility to become a diagnostic tool for neuropathy.

Limitations

The PTT technique indirectly measures the membrane excitability in a similar way as established methods of threshold tracking of the compound action potential. The main disadvantages with the perception threshold assessment are the influence of the central nervous system. To reduce the influence of the central nervous system a low frequency as well as low intensity stimulation are used during PTT since high frequencies and high intensities are prone to induce plasticity changes in the central nervous system (Klein, et al., 2004; Xia, et al., 2016). The variability between different subjects is of course substantial, but has been controlled for when using the data to fit the computational model by normalizing the data of the individual participants. The major cause of between subject variation in healthy participants is probably the distance from the cathode to the nerve fiber ending. Therefore, skin thickness and electrode placement may contribute most to the between subject variation. A major source of within subject variation is habituation which will cause the perception threshold to increase during the experiment. To reduce this effect, in the current study, the different durations of the ramp pulse were delivered in a randomized order.

| One of the limitations of the computational model is that it did not include any calcium |
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| channels which play an important role in regulating the excitability changes in nerve fibers. |
| Finally, the model did not include the influence of the electrical properties of the skin, but |
| assumed that this would not affect the activation of the nerve fibers. This is a simplification |
| as the capacitive properties of the tissue will smoothen the electrical field and thus the |
| electrical field at the nerve fibers will not be exactly the same as seen on the skin surface. An |
| additional limitation of the computational model is the number of unknown parameters, |
| particularly the maximum conductance of the voltage-gated ion channels. |
| |
| Author Contributions |

Conceptualization of the study was done by JT, TNH, KSF and CDM. All experimental data were collected and analyzed by TNH, DB, MV, and CDM. The computational work was performed by JT. The manuscript draft was written by JT and TNH. All authors revised the manuscript and approved the final version for publication.

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| 746 | |
| 747 | Figures |
| 748 | Figure 1. Study design. A. The definition of accommodation used in the current study. For |
| 749 | long ramp pulses (> 20 ms), reduced excitability may occur, and the intensity needed for the |
| 750 | subject to perceive the stimulus is increased (dashed line). B. The PTT technique. A surface |
| 751 | electrode was placed on the skin, and the subject reported if the stimulus was perceived. The |
| 752 | stimulus was a linear increased ramp current with the duration 1 ms, 10 ms, 25 ms, 50 ms, |
| 753 | 100 ms and 200 ms. C. The computational model. The model consists of two nerve fiber |
| 754 | models — one unmyelinated axon model representing the unmyelinated intraepidermal part |
| 755 | of an $A\delta$ fiber and one myelinated fiber model $(Aeta)$ representing a large fiber. The |
| 756 | extracellular potential was altered at the tip of the fibers to simulate the activation of nerves |
| 757 | by the electrodes. The morphology of the A eta model consists of three parts: node of Ranvier, |
| 758 | juxtaparanode and internode. The figures are not drawn to scale. |
| 759 | |
| 760 | Figure 2. Accommodation of cutaneous nerves. The perception threshold for different |
| 761 | duration of the ramp pulses for the pin electrode (A) and patch electrode (B.) C. Normalized |
| 762 | perception threshold. The error bars represent the standard error. The asterisk represents |
| 763 | statistically significant (* $p < 0.05$) different perception thresholds. |

| 764 | Figure 3. Voltage-gated ion channel currents during an action potential. The figures to the |
|-----|--|
| 765 | left represent an action potential generated in the $A\delta$ model and to the right the $A\beta$ model. |
| 766 | The extracellular potential was altered for 1 ms with the shape of a ramp pulse. The onset of |
| 767 | the stimulus occurred at time zero. A. Membrane potential. B. The small voltage-gated ion |
| 768 | channel currents. C. The large voltage-gated ion channel currents. The current density for |
| 769 | the potassium current was low for the K_{Dr} current (A $oldsymbol{eta}$ model) since the combined area of the |
| 770 | juxtaparanode was 3.33 times larger than the node of Ranvier. |
| 771 | |
| 772 | Figure 4. Accommodation generated by the computational model. A. The extracellular |
| 773 | potential alteration needed to generate an action potential which propagates to the end of the |
| 774 | nerve fiber model. B. The extracellular potential normalized to the 1 ms duration of the |
| 775 | stimulus. The generation of an action potential for different durations of the ramp stimulation |
| 776 | for the $A\delta$ model (C) and $A\beta$ model (D). The membrane potential for $A\delta$ (E) and $A\beta$ model |
| 777 | (F) when the extracellular potential alteration was increased from $10~\mathrm{mV}$ to $30~\mathrm{mV}$ |
| 778 | (duration=200 ms). The onset of the stimulus occurred at time zero. |
| 779 | |
| 780 | Figure 5. The influence of the maximum conductance on activation threshold. All currents |
| 781 | were increased by 30 %. For the larger currents, spiking sodium and delayed rectifier, the |
| 782 | maximum conductance was altered by 10% and 20% respectively. The green dotted line in |
| 783 | figure B represents the normalized extracellular potential when the maximum conductance of |
| 784 | both the K_M and delayed rectifier (K_{Dr}) currents was reduced by 50%. |
| 785 | |
| 786 | Figure 6. The influence of voltage-gated ion channels on accommodation. To study the |
| 787 | influence of specific different ion channels, either the conductances were altered, or the ion |
| 788 | channel was removed from the model. The maximum conductance of the K_{Dr} was varied in |

the two nerve fiber models (blue lines). All slow currents (K_M , Na_P , and HCN) were removed from the models (light blue line), and all the ion channels were removed from the model except the Na_{TTXs} for $A\beta$ model and Na_{TTXr} for the $A\delta$ model (light blue lines). To compensate for the high excitability, the conductance of the sodium channels was reduced by 54% for the case when all the ion channels were removed except for the spiking sodium channel. Note that accommodation could be generated in the $A\beta$ model when the model only Na_{TTXs} channel was implemented (light blue lines).

Figure 7. The influence of Inactivation of sodium channels. A. The membrane potential generated by a 10 mV ramp stimulation alteration of the extracellular potential (duration=200 ms). B. The total inactivation of the sodium channels (Na_{TTXs} or Na_{TTXr}) for all of the inactivation gates. C-F. The onset of the stimulus occurred at time zero. All the steady-state inactivation curves of either the Na_{TTXs} ($A\beta$ model) or Na_{TTXr} ($A\delta$ model) were shifted, and the accommodation recalculated. A negative shift is defined as a hyperpolarized shift of all of the steady-state inactivation curves and a positive shift as a depolarizing shift. To compensate for the general excitability, the maximum conductance of the Na_{TTXs} was increased ($A\beta$ model) or the Na_{TTXr} altered ($A\delta$ model) to retain a similar activation threshold (within 5%) for 1 ms ramp as in the control model (no shifts of the steady-state inactivation curves).













