

Temperature increases by kilohertz frequency spinal cord stimulation

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ABSTRACT

Introduction: Kilohertz frequency spinal cord stimulation (kHz-SCS) deposits significantly more power in tissue compared to SCS at conventional frequencies, reflecting increased duty cycle (pulse compression). We hypothesize kHz-SCS increases local tissue temperature by joule heat, which may influence the clinical outcomes.

Methods: To establish the role of tissue heating in kHz-SCS, a decisive first step is to characterize the range of temperature changes expected during conventional and kHz-SCS protocols. Fiber optic probes quantified temperature increases around an experimental SCS lead in a bath phantom. These data were used to verify a SCS lead heat-transfer model based on joule heat. Temperature increases were then predicted in a seven-compartment (soft tissue, vertebral bone, fat, intervertebral disc, meninges, spinal cord with nerve roots) geometric human spinal cord model under varied parameterization.

Results: The experimentally constrained bio-heat model shows SCS waveform power (waveform RMS) determines tissue heating at the spinal cord and surrounding tissues. For example, we predict temperature increased at dorsal spinal cord of 0.18–1.72 °C during 3.5 mA peak 10 KHz stimulation with a 40–10–40 μs biphasic pulse pattern, 0.09–0.22 °C during 3.5 mA 1 KHz 100–100–100 μs stimulation, and less than 0.05 °C during 3.5 mA 50 Hz 200–100–200 μs stimulation. Notably, peak heating of the spinal cord and other tissues increases superlinearly with stimulation power and so are especially sensitive to incremental changes in SCS pulse amplitude or frequency (with associated pulse compression). Further supporting distinct SCS intervention strategies based on heating; the spatial profile of temperature changes is more uniform compared to electric fields, which suggests less sensitivity to lead position.

Conclusions: Tissue heating may impact short and long-term outcomes of kHz-SCS, and even as an adjunct mechanism, suggests distinct strategies for lead position and programming optimization.

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Introduction

The emergence of kilohertz frequency (1–10 KHz) spinal cord stimulation (kHz-SCS) [1–7] for the treatment of neuropathic pain has engendered studies on new mechanisms of actions (MoA) [5,8–11]. Divergent clinical observations for conventional rate SCS and kHz-SCS suggest difference in MoA which could in turn inform distinct programming optimization strategies. Notably, kHz-SCS can provide an analgesic and side-effects profile distinct from conventional frequency (~100 Hz) SCS [11,12]. For example, kHz-

SCS does not produce the paresthesia associated with dorsal column activation in conventional SCS, and recent studies seemingly rule out direct activation of dorsal column fibers as the primary mechanism of action of kHz-SCS pain relief [13,14]. Wash-in time for the therapeutic benefit of conventional rate SCS is on the order of minutes, while responses to wash-in over a longer period [4]. Further indicating distinct MoA, kHz-SCS waveforms involve simultaneous decrease in pulse duration (well below membrane time constants) and increase in pulse frequency (beyond axon refractory periods) that challenge conventional models of stimulation [15,16].

Evidence against traditional neural MoA warrants investigation of other phenomena. We note that since the decrease in interpulse-interval (e.g. from 10 ms at 0.1 KHz to 0.1 ms at 10 KHz) is more drastic than the decrease in pulse duration (e.g. from 100 μs per

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phase at 0.1 KHz to 40 μ S per phase at 10 KHz [5,9]), kHz stimulation is associated with higher duty cycle – and the RMS power of a rectangular waveform varies positively with the square root of its duty cycle. Through the principle of joule heating, the power of current flow from an implanted lead can produce temperature increases around the lead [10,17–23]. Thus, kHz stimulation deposits more power in the tissue than conventional spinal cord stimulation and is therefore more likely to significantly heat the tissue immediately surrounding the stimulation site. A temperature increase and resultant thermal conduction into the spinal cord can, in turn, affect neuronal function [23] (e.g., via alteration of ion channel or neurotransmitters dynamics) and related biological functions (e.g., via vasodilation [24], heat shock protein expression [25]) depending on the degree of change. Tissue heating further encourages the expression of anti-inflammatory agents, such as heat shock proteins [26], over a period of time consistent with the extended wash-in times of kHz-SCS treatment.

Any form of electrical stimulation produces passive heating and the extent of induced temperature increases are specific to both the stimulation and local tissue properties, with various stimulation and environmental parameters affecting the degree to which heating occurs [19,23,27]. Key stimulation parameters are the stimulation waveform (based on stimulator programming) and electrode montage (based on lead placement), which together with tissue anatomy and electrical conductivity determine joule heat deposition. An implanted stimulator is a constant energy source which will produce unlimited temperature increases without passive (e.g. heat conduction by CSF) or active (e.g. spinal tissue blood perfusion) heat dissipation by the tissue. As such, heating analysis depend on tissue properties such as thermal conductivity, metabolic rate, and blood perfusion; not only of the stimulation target but surrounding tissues. Indeed, we postulate that the local environment around SCS leads is especially conducive to temperature increases, namely the low conductivity of fat and enclosed anatomy of the vertebral canal. If heating due to these factors is sufficient during kHz frequency SCS to shape beneficial responses, then joule heating by SCS may be an adjuvant mechanism underlying therapy. However, the degree of heating during kHz-SCS, including as aggravated by increased power deposition due to pulse compression and/or the enclosed spinal environment, remains unexplored.

The objective of this study was to assess, for the first time, whether an increased duty-cycle (and so power) of High-Rate spinal cord stimulation will produce significant temperature increases in the spinal cord. Prior experimental and modeling studies of conventional non-invasive and invasive forms of brain stimulation has suggested minimal heating under normal device operation (less than 1 °C) [18,28–31]. This study predicts the degree of tissue temperature rises driven by SCS joule heat, and characterizes the role of SCS waveform (including frequency, pulse width, and amplitude) and tissue properties. We measured temperature increases around an experimental SCS lead in a bath to verify a finite-element-model of SCS joule heat. We confirmed the dependence of temperature rise only on the power of the stimulation waveform, independent of other parameters. Finally, we predicted temperature increases during conventional and kHz-SCS at the dorsal spinal cord under passive and active bio-heat conditions in a geometric human spinal cord FEM model.

Method

Bath phantom study

Saline bath phantom

Thermal and electrical conductivity measurements taken to verify the general heat transfer model were performed in a

cylindrical glass container (diameter: 90 mm and height: 130 mm) with three varied NaCl concentrations (154 mmol/L, 34.22 mmol/L, and 3.42 mmol/L (approximating cerebrospinal fluid, meninges, and epidural space respectively). A thermal conductivity meter (Therm Test Inc., Canada) and an electrical conductivity meter (Jenco Instruments, Inc., San Diego, CA) measured the thermal and electrical properties of the saline solutions at 37 °C (core spinal cord temperature approximation). The measured corresponding conductivity values for each molar concentrations were: electrical conductivity (σ): 1.62 S/m, 0.47 S/m, and 0.047 S/m; and thermal conductivity (κ): 0.6268 W/(m.K), 0.6317 W/(m.K), and 0.6319 W/(m.K) respectively.

In vitro stimulation

For the saline bath experiments, an experimental polyurethane SCS lead with 4 Platinum/Iridium electrode contacts (1.35 mm electrode diameter, 3 mm electrode length, 1 mm inter-electrode spacing) was placed at the center of the cylindrical container. The cylindrical container was then immersed in a temperature-controlled water bath (280 x 160 x 150 mm³) maintained at ~37 °C (Fig. 1A) and baseline temperature was stabilized for > 60 min. Three different waveforms, namely sinusoidal, square, and a symmetric charge-balanced biphasic pulse waveforms mimicking the characteristics and parameters of clinical SCS waveforms (described by leading pulse duration, inter-pulse interval, recovery pulse duration), were generated using a function generator (AFG320, Tektronix, Beaverton, OR, USA). The generated waveforms were passed through a custom designed high-bandwidth linear current isolator to the experimental SCS lead. (Distal) Electrode contact 1(E1) and (proximal contact) 4 (E4) of the experimental SCS lead were energized for all saline bath experiments. Tested stimulation intensities were 1–7 mA (peak) using rates of 0.1 KHz to 10 KHz. Only for phantom verification, biphasic rectangular waveform pulse widths of each phase (40 μ s) and interphases (10 μ s) were kept constant such that the duty cycle increased directly with stimulation frequency.

Temperature measurement and analysis

A fiber optic temperature probe (STS Probe Kit, LumaSense Technologies, Inc. CA, USA) sensed by a fiber optic thermometer (± 0.1 °C accuracy at calibration temperature, m600 FOT LAB KIT, LumaSense Technology, CA, USA) was positioned in the proximity of E4 to measure temperature increases during stimulation (Fig. 1A). We measured the peak temperature change in the bath radially from E4 (1 mm, 2 mm, 3 mm, and 4 mm) during stimulation as a function of peak stimulation amplitudes (1–7 mA), over a range of stimulation frequencies (0.1 KHz, 1 KHz, 5 KHz, 10 KHz, and 20 KHz) for sinusoidal, square, and SCS pulsed waveforms. Measured temperature was digitized using TrueTemp data acquisition and graphing software (60 samples/measurement and 1 s measurement interval, LumaSense Technologies, Inc. CA, USA). Temperature was normalized with respect to the initial temperature (~37 °C), which was considered baseline.

Computational models and solution method

Bioheat model of spinal cord

Human spinal cord was simulated as a computer-aided design (CAD) derived model comprising seven compartments namely vertebrae (lower thoracic region, T8–T11), intervertebral disc, surrounding soft-tissues (minimally perfused), epidural fat, meninges, cerebrospinal fluid, and spinal cord (white matter and grey matter combined; Fig. 2). The dimensions of the individual tissues, modelled as isotropic homogenous volume conductors, were based

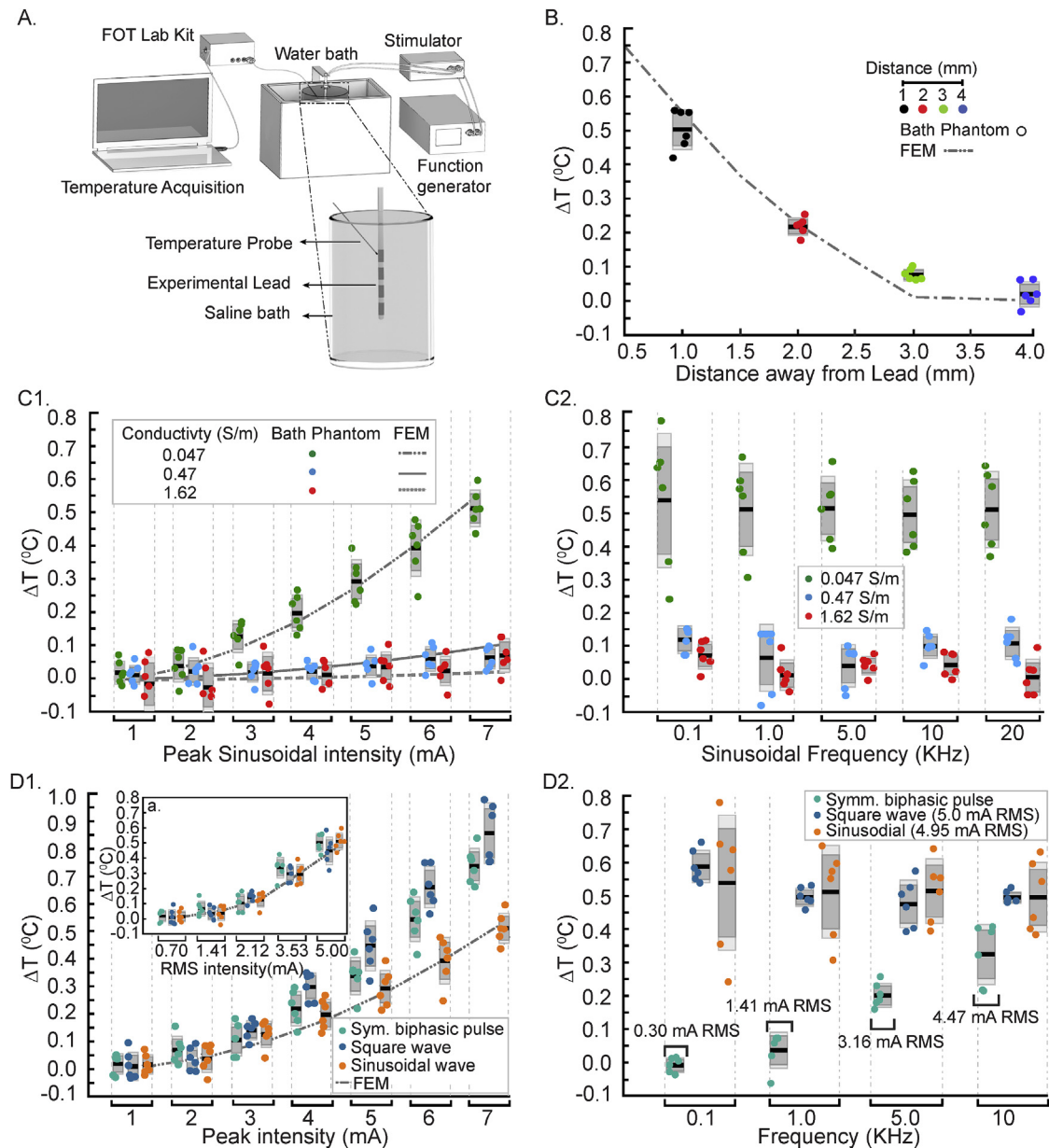


Fig. 1. Measurement of temperature increases in phantom preparation across rates, waveforms (SCS, sinusoidal, square) and conductivities and verification of FEM SCS heat model. Dark grey box: standard error of the mean; Light grey box: standard deviation; black line: mean of the data, and the dots are the individual ΔT measurements. (A) Schematic of a salt bath experimental set up with an experimental SCS lead in a salt bath heated in a water bath, function generator driving custom isolator energizing to the experimental SCS lead, and optical temperature probe mounted on a micro-manipulator. (B) ΔT measured at a radial direction away from the electrode contact (E4, positive polarity) when applying a 10 KHz Symmetric biphasic pulsed waveform at 5 mA peak in a low conductivity saline bath, and the corresponding FEM heat model. Spatial temperature field decreased with increasing radial distance as predicted with the highest temperature increases (mean $\sim 0.5^\circ\text{C}$) measured proximal to the lead (1 mm). (C1) For 10 KHz sinusoidal waveform, temperature increases as a function of peak stimulation intensity in varied saline bath conductivities: Green: 0.047 S m^{-1} ; Blue: 0.47 S m^{-1} ; Red: 1.62 S m^{-1} . Experimental (dots) and predicted (broken lines) ΔT significantly increased with stimulation intensity with higher sensitivity in lower conductivity saline baths. (C2) For 10 KHz sinusoidal waveform, ΔT measurement at varied stimulation frequencies (0.1, 1, 5, 10, 20 kHz) and conductivities. Temperature increases were independent of frequency and consistently higher for low saline conductivity ($p < 0.01$). (D1) ΔT across different waveforms and stimulation intensities (Main Panel: Peak; Inset: RMS matched) with frequency fixed at 10 KHz. (D2) ΔT s for the different waveforms across various frequencies at RMS stimulation intensities (Square, 5 mA; Sinusoidal, 4.95 mA; Symmetric biphasic pulsed, frequency-dependent RMS). (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

on human cadaveric spinal cord from prior studies [32]. The diameter of spinal cord with dorsal roots was fixed (spinal cord, 6.4 mm; dorsal roots, 0.5 mm) and the thickness of the adjacent tissues were: CSF, 2.0 mm; meninges, 0.5 mm; and epidural fat 1.0 mm. We note that in situ, the diameter of the spinal cord varies along the vertebral column. Two SCS clinical leads were modelled and placed epidurally in a minimally staggered bilateral fashion (SCS Lead 1, 1 mm distal to the mediolateral midline at T8; SCS Lead 2, 0.5 mm away from SCS Lead 1 and proximal to the mediolateral

midline at T9; Fig. 2A2). We energized only the first SCS lead; the second lead was passive, positioned to mimic a clinical placement, and used to assess the impact of the presence of a passive lead on heat dispersion [33]. The finite element method (FEM) model was solved using Pennes' bioheat equation governing joule heating during electrical stimulation (Laplace equation for electrostatics ($\nabla(\sigma\nabla V) = 0$ where V is potential and σ is conductivity), metabolic heat generation rate (Q_{met} or MHG), and blood perfusion rate (ω_b or BPer) in the tissues as mentioned below:

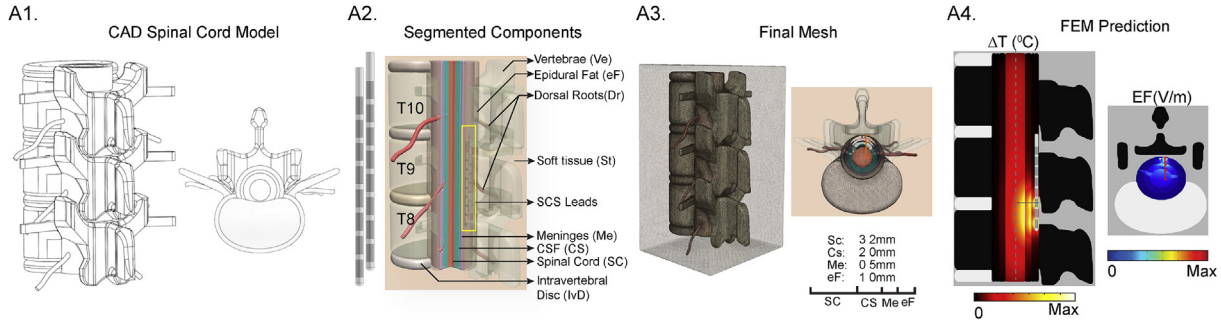


Fig. 2. FEM SCS heat transfer model architecture and work flow. (A1) CAD derived spinal cord model with dimensions based on human cadaver study [68]. (A2) Geometry of sample clinical human SCS lead and a placement of the lead in a segmented spinal cord with dorsal roots and surrounding tissues. SCS lead was positioned into the epidural space between lower thoracic (T8–T10) vertebral region. Two leads were modelled as to mimic clinical intervention, positioned in a minimally staggered bilateral fashion. Epidural space was modelled with fatty tissues. (A3) Resulting volumetric mesh of spinal tissues. (A4) The numerical solution predicted ΔT ($^{\circ}\text{C}$) and EF (V/m) in the spinal tissues. Black line in the temperature profile plot and red line in the EF plot represent radial direction from the lead where temperature was sampled. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

$$\rho C_p \nabla T = \nabla \cdot (\kappa \nabla T) - \rho_b C_b \omega_b (T - T_b) + Q_{met} + \sigma |\nabla V|^2 \quad (1)$$

where ρ , C_p , T , σ , and κ represent tissue density, specific heat, temperature, electrical conductivity, and thermal conductivity respectively. Biological properties of blood such as density (ρ_b), specific heat (C_b), and temperature (T_b) were assumed constant in all vascular spinal tissues (vertebrae, meninges, spinal cord) and the corresponding values were 1057 kg/m^3 , 3600 J/(kg.K) , and 36.7°C respectively. Blood perfusion rate (ω_b) values were tissue specific and were in the range of $0.0003\text{--}0.008 \text{ s}^{-1}$ [34,35]. In spinal tissues, metabolic activities due to local spinal cord metabolism and enhanced metabolism in response to SCS generates thermal energy [36]. Blood circulation also plays a significant role in transporting thermal energy across the spinal tissues through convection [37]. We considered the blood temperature in the spinal tissues to be 0.3°C less than core spinal cord temperature (37°C). We investigated how the interaction between metabolic heat generation and blood perfusion modulates kHz-SCS induced temperature increases. Prior to the application of kHz-SCS, the metabolic heat generation rate required to balance the initial spinal cord temperature was calculated using equation (2) [37,38] for the aforementioned perfusion rates as:

$$Q_{met} = \rho_b C_b \omega_b (T - T_b) \quad (2)$$

where T and T_b are initial spinal cord and blood temperature. The calculated Metabolic Heat Generation (MHG) and the corresponding Blood Perfusion (BPer) values were given as; spinal cord and meninges ($Q_{met}, 9132 \text{ Wm}^{-3}$; $\omega_b, 0.008 \text{ s}^{-1}$), vertebrae ($Q_{met}, 342 \text{ Wm}^{-3}$; $\omega_b, 0.0003 \text{ s}^{-1}$), and minimally perfused soft-tissues ($Q_{met}, 457 \text{ Wm}^{-3}$; $\omega_b, 0.0004 \text{ s}^{-1}$). The balanced Q_{met} values approximated prior experimental measurements [34,35,39]. CSF convection is not incorporated, but could be considered in future bio-heat models.

Mimicking clinical montages and waveforms, we energized electrode contacts E1 and E3 of the clinical SCS Lead 1 in a bipolar configuration (8 mm center-to-center electrode distance). Maximum temperature increases by conventional and kHz-SCS using rectangular waveforms for varied peak amplitudes (1, 2, 3, 3.5, 4, 5 mA), frequencies and pulse widths (50 Hz (200 μs), 100 Hz (200 μs), 1 KHz (40 μs and 100 μs), 5 KHz (40 μs), and 10 KHz (40 μs) were predicted and compared between active (bioheat) and passive heating cases at three different locations namely, at the distal edge E3 of the clinical SCS Lead 1 ($\sim 0.01 \text{ mm}$ from the surface of the

lead), at the proximal surface of the dorsal root to the SCS lead, and at the surface of spinal cord ($\sim 3.5 \text{ mm}$ radial from the E3 electrode).

Boundary and initial condition

To model each stimulation waveform, we applied corresponding static RMS values (see phantom and model Results for justification). The accuracy of RMS intensities calculated analytically for a given intensity, frequency, and pulse width (see equation (3)) were confirmed experimentally by stimulation across a resistive load (1 K Ω) with voltage acquisition using a digital mixed signal oscilloscope (MSO2024, Tektronix, OR, USA, $\pm (100 \text{ mv} + 3\%$ of threshold)), a DAQ (NI PCI 5922, National Instruments, TX, USA, $\pm 500 \text{ ppm}$ (0.05%) of input + 50 μV), and a digital multimeter (DMM 7510 7 $\frac{1}{2}$ Digit Graphical Sampling Multimeter, Tektronix, OR, USA, $\pm 60 \text{ ppm}$ 0.0014% of input). The error in calculated versus measured RMS values was less than 5%.

$$\begin{aligned} I_{RMS} &= \sqrt{\frac{1}{T} \int_0^t I(t)^2_{peak} dt} \\ &= I(t)_{peak} \sqrt{\frac{t}{T}} \\ &= I_{peak} \sqrt{D} \end{aligned} \quad (3)$$

where I_{peak} is the peak bipolar stimulation intensity, I_{RMS} is the corresponding RMS value, T is the pulse duration, t is the pulse width, and D is the duty cycle.

A static inward normal current density ($J_{norm, RMS}$) corresponding to the stimulation current intensity (I_{RMS} , Table 1) was injected through E1, and E3 was set as the return (producing a bipolar configuration). The electrical and thermal conductivities of the electrode contacts and the inter-electrode spacing were $4 \times 10^6 \text{ S/m}$ and 31 W/(m.K) , and $\sigma = 1 \times 10^{-15} \text{ S/m}$; $\kappa = 0.0262 \text{ W/(m.K)}$ respectively [36]. The outer boundaries of the spinal cord and the surrounding tissues were considered electrically insulated.

For the thermal boundary conditions, the temperature at the outer boundaries of the spinal column was fixed at core body temperature (37°C) [35,39] with an assumption of no convective heat loss to the ambient temperature, no convective gradients across spinal surrounding tissues, and no SCS-induced heating at the model boundaries [34]. The initial temperature of the tissues was assumed to be 37°C , and thermo-electrical properties of biological tissues were based on average literature values [40,41]. Intravertebral disc ($\sigma = 0.830 \text{ S/m}$; $\kappa = 0.49 \text{ W/(m.K)}$) and csf

Table 1
FEM SCS Bio-heat model predicts temperature increases (ΔT) in fat around the Lead, the surface of spinal cord (SC), and the proximal origin of dorsal Root, as a function of stimulation waveform and intensity (I), for passive (A.) and active (B.) tissue models. For each waveform (top row) the corresponding Pulse Compression Factor (PCF) is indicated (see Methods). For each waveform and intensity simulation, the corresponding RMS is indicated. In addition to increments of 1 mA peak, 3.5 mA peak is also simulation (grey rows).

A. Passive heating																								
PCF	10 KHz (40-10-40 μs)				5 KHz (40-10-40 μs)				1 KHz (40-100-40 μs)				1 KHz (100-100-100 μs)				100 Hz (200-100-200 μs)				50 Hz (200-100-200 μs)			
	6.32				4.47				2.00				3.16				1.41				1.00			
I (mA)	ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)			
Peak	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root
1.0	0.89	0.34	0.07	0.03	0.63	0.17	0.04	0.02	0.28	0.03	0.01	0.00	0.47	0.08	0.02	0.01	0.20	0.02	0.00	0.00	0.14	0.01	0.00	0.00
2.0	1.78	1.32	0.29	0.12	1.26	0.67	0.14	0.06	0.56	0.13	0.03	0.01	0.89	0.33	0.07	0.03	0.40	0.07	0.01	0.01	0.28	0.03	0.01	0.00
3.0	2.68	3.03	0.65	0.27	1.89	1.49	0.32	0.14	0.84	0.29	0.06	0.03	1.34	0.75	0.16	0.07	0.60	0.15	0.03	0.01	0.42	0.08	0.02	0.01
3.5	3.13	4.11	0.84	0.35	2.21	2.05	0.43	0.18	0.98	0.40	0.09	0.04	1.57	1.01	0.22	0.09	0.70	0.21	0.04	0.02	0.49	0.10	0.02	0.01
4.0	3.57	5.24	1.15	0.49	2.53	2.63	0.57	0.24	1.12	0.53	0.11	0.05	1.79	1.31	0.28	0.12	0.80	0.27	0.06	0.02	0.57	0.13	0.03	0.01
5.0	4.47	8.31	1.79	0.76	3.16	4.15	0.90	0.38	1.41	0.82	0.18	0.07	2.24	2.04	0.44	0.19	1.00	0.41	0.09	0.04	0.71	0.21	0.04	0.02
B. Active heating (Perfusion and MHR)																								
PCF	10 KHz (40-10-40 μs)				5 KHz (40-10-40 μs)				1 KHz (40-100-40 μs)				1 KHz (100-100-100 μs)				100 Hz (200-100-200 μs)				50 Hz (200-100-200 μs)			
	6.32				4.47				3.16				2.00				1.41				1.00			
I (mA)	ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)				ΔT (°C)			
Peak	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root	RMS	Lead	SC	Root
1.0	0.89	0.30	0.02	0.01	0.63	0.15	0.01	0.00	0.28	0.03	0.00	0.00	0.47	0.08	0.00	0.00	0.20	0.02	0.00	0.00	0.14	0.01	0.00	0.00
2.0	1.78	1.19	0.12	0.03	1.26	0.59	0.05	0.01	0.56	0.12	0.01	0.00	0.89	0.31	0.02	0.01	0.40	0.06	0.00	0.00	0.28	0.03	0.00	0.00
3.0	2.68	2.69	0.27	0.06	1.89	1.33	0.13	0.03	0.84	0.26	0.02	0.01	1.34	0.68	0.07	0.02	0.60	0.13	0.01	0.01	0.42	0.07	0.00	0.00
3.5	3.13	3.60	0.37	0.08	2.21	1.78	0.18	0.04	0.98	0.35	0.03	0.01	1.57	0.92	0.09	0.02	0.70	0.19	0.01	0.01	0.49	0.09	0.00	0.01
4.0	3.57	4.71	0.49	0.10	2.53	2.37	0.24	0.05	1.12	0.45	0.04	0.01	1.79	1.21	0.12	0.03	0.80	0.24	0.02	0.01	0.57	0.12	0.01	0.01
5.0	4.47	7.25	0.77	0.16	3.16	3.69	0.38	0.08	1.41	0.73	0.07	0.02	2.24	1.89	0.19	0.04	1.00	0.39	0.03	0.01	0.71	0.19	0.01	0.01

* Pulse Compression factor (PCF).

($\sigma = 1.65$ S/m; $\kappa = 0.57$ W/(m.K) are avascular, and therefore have no BPer and MHG, whereas the other remaining tissues are vascularized and have BPer and MHG as listed: epidural fat ($\sigma = 0.025$ S/m; $\kappa = 0.21$ W/(m.K), $\omega_b = 0.0001$ s $^{-1}$, $Q_{met} = 58$ Wm $^{-3}$) soft tissues ($\sigma = 0.15$ S/m; $\kappa = 0.47$ W/(m.K), $\omega_b = 0.0004$ s $^{-1}$, $Q_{met} = 457$ Wm $^{-3}$), vertebrae ($\sigma = 0.01$ S/m; $\kappa = 0.32$ W/(m.K), $\omega_b = 0.0003$ s $^{-1}$, $Q_{met} = 342$ Wm $^{-3}$), meninges ($\sigma = 0.368$ S/m; $\kappa = 0.44$ W/(m.K), $\omega_b = 0.008$ s $^{-1}$, $Q_{met} = 9132$ Wm $^{-3}$), and spinal cord ($\sigma = 0.126$ S/m; $\kappa = 0.51$ W/(m.K), $\omega_b = 0.008$ s $^{-1}$, $Q_{met} = 9132$ Wm $^{-3}$). When indicated, these “standard” tissue values were manipulated by either 1) doubling or halving the electrical and/or thermal conductivities of a given compartment, or 2) by substituting properties across compartments.

Saline bath phantom FEM

SCS saline bath phantom was modelled using equation (1) while eliminating the biological tissue parameters. The FEM Phantom model was parameterized based on the dimensions, conductivity, and initial temperature of the experimental set-up. As tested, we simulated one SCS experimental lead centrally placed in a saline bath phantom. For the electrical boundary conditions, a uniform RMS current density was applied at E4 (anode) and return at E1(cathode). The outer boundaries of the bath were considered electrically insulated. For thermal boundary conditions, the external boundary temperature and the initial temperature of the bath were fixed at 37 $^{\circ}$ C. To account for the voltage drop due to the electrode-saline interface [42–44], a correction factor of 50% was applied in phantom stimulation [18].

Model construction and computational method

Human spinal cord and saline bath phantom models were CAD derived, assembled in SolidWorks 2016 (Dassault Systemes

Americas Corp., Ma, USA), and imported using Simpleware ScanIP (Synopsys Inc., CA, USA). The entire volume of the spinal tissue and the electrode assembly was 83.0 x 74 x 108 mm 3 . Prior to the segmentation, tissues were resampled to have an isotropic resolution of 0.2 mm 3 . Resampled images were segmented into seven tissues compartments along with the T8-T11 positioned SCS lead assembly using a combination of automatic and manual segmentation filters (Fig. 2A2). Using a voxel-based meshing algorithms of ScanIP, an adaptive tetrahedral mesh was generated. The final model size resulting from multiple mesh densities refinement contained approximately 4,600,000 tetrahedral elements for the full anatomy of spinal cord model and approximately 320,000 tetrahedral elements for the saline bath model (Fig. 2A3). The meshes were imported into COMSOL Multiphysics 5.1 (COMSOL Inc., MA, USA) to computationally solve the FEM model. The SCS model was solved for both passive heating (joule heating, without BPer and MHG) and active heating (bioheat, with BPer and MHG) conditions. The baseline temperature gradient for the active heating case was predicted by first solving the heat transfer model in the absence of electrical stimulation. In passive heating, the baseline temperature gradient was set to zero. The Saline bath model was solved only for passive heating condition. Both phantom and SCS models were solved under steady state assumption and corresponding temperature increases and field intensities were quantified. Heat flux and field intensity streamlines (seeded at selected tissue boundaries and proportional in diameter to the logarithm of corresponding magnitudes) were plotted to illustrate the overall distribution across tissues (Fig. 3).

Statistics and analysis

Normality test on temperature increases were conducted using Lilliefors corrected K-S test statistical test. A two-way repeated

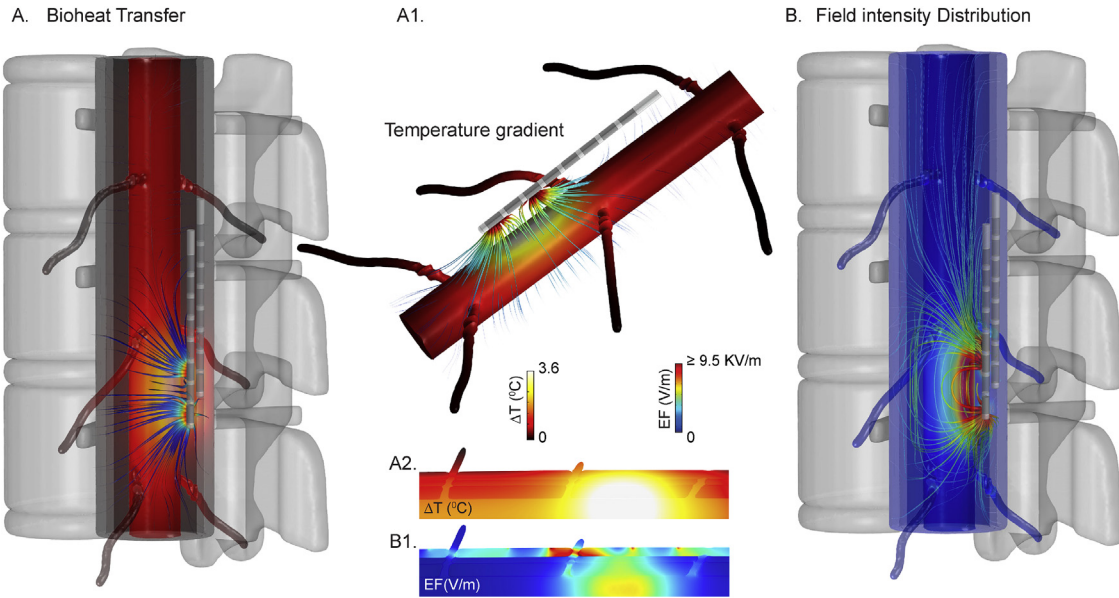


Fig. 3. FEM Bioheat transfer model of Spinal Cord Stimulation predicts temperature changes and electric field intensities. (A) Temperature increases at the spinal cord modelled with dorsal roots, and surrounding tissues for a metabolic heat and blood perfusion modulated high rate spinal cord stimulation (kHz-SCS). Temperature gradient streamlines originated from SCS leads to the spinal cord and dorsal roots are depicted. (A1) Accumulated heat (shown as heat flux streamlines) transported throughout spinal cord and dorsal roots during kHz-SCS. (A2, B2) illustrates predicted temperature and electric field (EF) spatial distribution on a spinal cord, dorsal roots, and other adjacent tissues. (B) Field intensity distribution across spinal cord and surrounding tissues. EF strength across each tissue are depicted by electric field streamlines seeded on every tissue boundary.

measure analysis of variance (ANOVA) was used to access the statistical differences in ΔT across different tested conditions (stimulation intensity, waveforms, frequencies, conductivities). A critical value (p) < 0.01 was accepted as a statistical difference between the groups. Further significance between groups were verified using Post hoc Scheffe's test (corrected multiple comparisons). The statistical relations between the experimental data the FEM data was evaluated through a simple linear regression MATLAB (R2016a, MathWorks, MA, USA).

The standard forms of power law tested super-linearity between the RMS and temperature increases, using a linear least squares fitting technique derived by Gauss and Legendre with a power function [45] given as:

$$\Delta T = A * RMS^\beta \quad (4)$$

where ' β ' is the power, and 'A' is the proportionality constant. The value of ' β ' determines the category of the relationship ($\beta = 1$, linear; $\beta > 1$, super-linear; $\beta < 1$, sublinear). Formulating the power function further on a log-log scale yields:

$$\ln(\Delta T) = \ln(A) + \beta * \ln(RMS) \quad (5)$$

Equation (5) is a straight line with a slope ' β ' and a y-intercept of $\ln(A)$. Linear least square fit of the logarithmic data yields the correlation (r^2) [45].

Pulse Compression Factor per stimulation intensity (PCF) captures the increase in RMS of a High-Rate waveform ($RMS_{High-Rate}$) compared to a conventional 1 mA peak 50 Hz 200 μ s pulse-width waveform (RMS_{50}):

$$RMS_{High-Rate} = I_{peak} * PCF * RMS_{50} \quad (6)$$

$$PCF = 10 * \sqrt{Pw * f} \quad (7)$$

where 'Pw' and 'f' are pulse width (sec) and frequency (Hz) for a given High-Rate waveform.

Results

Phantom measurements and model verification

A specially designed chamber was used to quantify temperature increases around an experimental SCS lead in a saline bath using varied waveforms (Fig. 1A). A micro-manipulator mounted optical temperature probe mapped steady-state temperature increases during stimulation with varied waveforms. As predicted by the FEM, temperature increases when applying a 10 KHz symmetric biphasic pulsed waveform at 5 mA peak intensity in a low conductivity saline phantom was maximal near energized electrodes and decreased with radial distance (Fig. 1B). In separate experiments, salt bath conductivity was varied by saline concentration. The main effect of saline bath conductivity and stimulation intensities (1–7 mA peak sinusoidal) was significant ($F(2, 105) = 218.95$, $p < 0.01$ and $F(6, 105) = 42.03$, $p < 0.01$, a two-way ANOVA). The interaction between these factors on ΔT was also significant; ($F(12, 105) = 19.88$, $p < 0.01$). Temperature increases were measured to be significantly greater in the lower saline bath conductivity (0.047 S/m) than in the other two saline bath conductivities (0.47 S/m and 1.62 S/m; Post-hoc Scheffe's test, Fig. 1C1). Across different saline conductivities at different sinusoidal frequencies, the measured temperature increases were significant; $F(2, 75) = 256.25$, $p < 0.01$. ΔT was higher at lower conductivity saline bath (Fig. 1C2).

Temperature increased by up to $\sim 1^\circ\text{C}$ with stimulation amplitude during stimulation using all 10 KHz waveforms (symmetric biphasic pulse, square, sinusoidal). In addition, when considering only peak intensities, higher ΔT was observed during stimulation using pulsed and square waveforms versus the sinusoidal waveform ($F(2, 105) = 41.14$, $p < 0.01$). However, this effect was found to be directly related to the RMS of the waveform (Fig. 1D1a) and not to the specific shape of the stimulation waveform ($F(2, 75) = 1.11$, $p > 0.01$).

In a separate series, temperature increases were measured across varied frequencies for all waveforms (symmetric biphasic

pulse, square, sinusoidal) in a low conductivity saline bath with 5 mA peak current intensity (corresponding RMS: sinusoidal waveform, 4.95 mA; square waveform, 5 mA; in pulsed waveform, RMS varies with frequency (Fig. 1D2). There was a main effect of stimulation waveforms on ΔT ; $F(2, 60) = 133.44$, $p < 0.01$. Temperature increases (0 – $\sim 0.4^\circ\text{C}$) across frequencies for symmetric biphasic pulsed waveform were significant ($p < 0.01$); however, for true square and sinusoidal waveforms, ΔT did not increase significantly across frequencies ($p > 0.01$). Temperature rises appeared to reflect the increase in duty cycle and RMS only for the symmetric biphasic pulsed waveform. Conversely, significantly higher temperatures were measured overall at the 5 mA peak intensity for sinusoid and square waveform compared to the pulsed waveform—reflecting the 100% duty cycles and therefore higher RMS values of the sinusoid and square waveforms.

Computational FEM predictions of the phantom using the experimental lead and waveforms were correlated with experimental temperature increases measurement at varied saline conductivities ($(R^2 = 0.24, F(1, 40) = 12.20, p < 0.01, 1.62 \text{ S/m}; R^2 = 0.26, F(1, 40) = 13.70, p < 0.01, 0.47 \text{ S/m}; R^2 = 0.84, F(1, 30) = 201.84, p < 0.01, 0.0047 \text{ S/m})$ (Fig. 1C1). Computationally predicted and measured temperature increases were strongly correlated across different RMS stimulation intensities ($R^2 = 0.86, F(1, 27) = 167.39, p < 0.01$ (Fig. 1D1a). Accordingly, a strong association between ΔT s were established along radial direction away from the experimental SCS lead; $R^2 = 0.96, F(1, 21) = 495.59, p < 0.01$ (Fig. 1B).

Computational model of heating by SCS: influence of waveform with standard tissue parameters

Using a FEM bio-heat computational models of human spinal cord stimulation, tissue temperature increases were predicted under varied stimulation parameters (Table 1) for passive heating and active conditions initially using “standard” tissue parameters (see Methods). Six representative SCS waveforms were simulated, with selected frequency and duty cycle (corresponding Pulse Compression Factor noted in table; see Discussion), each with varied peak intensity from 1 to 5 mA (corresponding resultant RMS noted in table). For each waveform and intensity, we tabulate the maximum ΔT around the SCS clinical lead (E3 contact), at the proximal surface of the dorsal root to the SCS lead ($\sim 1 \text{ mm}$ lateral to the stimulating lead), and at the surface of spinal cord ($\sim 3.5 \text{ mm}$ radial to the stimulating lead).

From this analysis, several important predictions emerge. Heating under the standard active model (which includes blood perfusion (BPer) and metabolic heat generation (MHG)) was lower than the standard passive model (where BPer and MHG were absent). Maximum temperature increases were generated around the SCS clinical lead (the epidural fat). Temperature increases were relatively higher for waveforms with a higher Pulse Compression Factor. Both active and passive heating increased with stimulation RMS, and so with intensity or Pulse Compression Factor, in a super-linear manner (e.g. doubling stimulation intensity or Pulse Compression Factor doubles RMS and results in a > 2 -fold increase in temperature (Fig. 4). While relative temperature increases were more sensitive to intensity than Pulse Compression Factor, the highest temperature increase were predicted under high Pulse Compression (e.g. the 10 KHz waveform). For example, using a conventional 50 Hz waveform (PCF: 1.0), temperature at the spinal cord (SC) increased $< 0.05^\circ\text{C}$ even at 5 mA peak (RMS: 0.71) while using a 10 KHz waveform (PCF: 6.32) temperature at the spinal cord (SC) increased $\sim 1^\circ\text{C}$ at 5 mA peak (RMS: 4.47).

Dependence of temperature increase on RMS (and so Intensity or Pulse Compression Factor) was modeled assuming a power law relationship, which results in a linear log-log dependence (see

Methods). Surprisingly, and despite the complexity of the standard tissue model, this fit sufficiently and reliably predicted temperature increases. Slope (β) approached 2 (i.e. temperature increasing with the square of RMS) – a super-linear ($\beta > 1$) sensitivity of temperature to RMS. The proportionality constant (A) increased across fat (Lead), Spinal Cord, and Root compartments, all relatively higher in the passive (Fig. 4A1) versus active (Fig. 4B1) tissue model.

Computational model of SCS: parameter sensitivity analysis with fixed waveforms

Living tissue possess complex thermo-electrical properties [45] and these properties are tissue specific. In the active model, we predicted the sensitivity of SCS temperature to tissue properties by halving or doubling the thermal and/or electrical conductivity (from the standard model; see Methods) of each tissue compartment. At 3.13 mA RMS (as for a 10 KHz SCS waveform with 3.5 mA peak), we considered a significant change in predicted temperature as $> 0.03^\circ\text{C}$ and $> 8\%$ from the standard model. No simulated changes in passive thermal and/or electrical conductivity at any tissue, except epidural fat (eF), produced a significant temperature change at the Lead, Spinal Cord, or Root. However, increases or decreases in epidural fat electrical conductivity significantly decreased or increased temperature across tissue compartments, respectively. The resulting predicted range of temperature increases using waveforms with 3.13 mA RMS were (Passive Model Range; Active Model Range): Lead (1.53 – 11.57°C ; 1.25 – 10.77°C), Spinal Cord (0.42 – 1.72°C ; 0.18 – 0.72°C), and Root (0.17 – 0.75°C ; 0.04 – 0.15°C).

We considered the sensitivity and fit of the power-law function across tissue properties, specifically varying fat electrical (σ) and thermal (k) conductivity (doubling and halving). In all tissue conditions, the linearity of log-temperature versus log-RMS confirmed a power-law fit, with consistently super-linear sensitivity ($\beta > 1$). Thus, for each tissue model, temperature could be predicted reliably by simply the corresponding power law function parameters, A and β (Fig. 4). In the passive model, β approached 2 across conditions. In the active model β could exceed 2, reflecting variance at low RMS, but not sensitivity at high RMS. The proportionality constant (A) varied more significantly across model parameters and tissue compartments, particularly near the Lead (Fig. 4A1, 4B1, 4B2).

To evaluate the contribution of peripheral spinal tissues on the temperature increases, we considered series of idealized models starting with uniform epidural fat and then sequentially adding adjacent tissues, under both active and passive model conditions (Fig. 5). The order of simulated tissues and predicted maximum temperature increases at locations corresponding to Lead position (“Lead”), Spinal Cord surface (“SC”), and dorsal Root surface (“Root”) are reported for both passive heating and active heating conditions (3.13 mA RMS at 10 KHz; Table 2). Maximum temperature increases and penetration (from the lead inward) is predicted in the uniform epidural fat model, with a relatively shallow electric field profile. The addition of Soft tissue (St), Vertebrae (Ve), and Intravertebral Disc (IVD) compartments and subsequent reduction of the size of the epidural fat layer(?) result in an incremental reduction in predicted temperatures increases – which is consistent with the notion that fat tissue properties are the most conducive to heating. The relative reduction in temperature between the active and passive models, as well as the reduction in electric field (which is always the same across active and passive models) emphasize these variables can change independently.

Further addition of Meninges (Me) to the model reduced predicted temperature rises notably in both relatively interior (Spinal

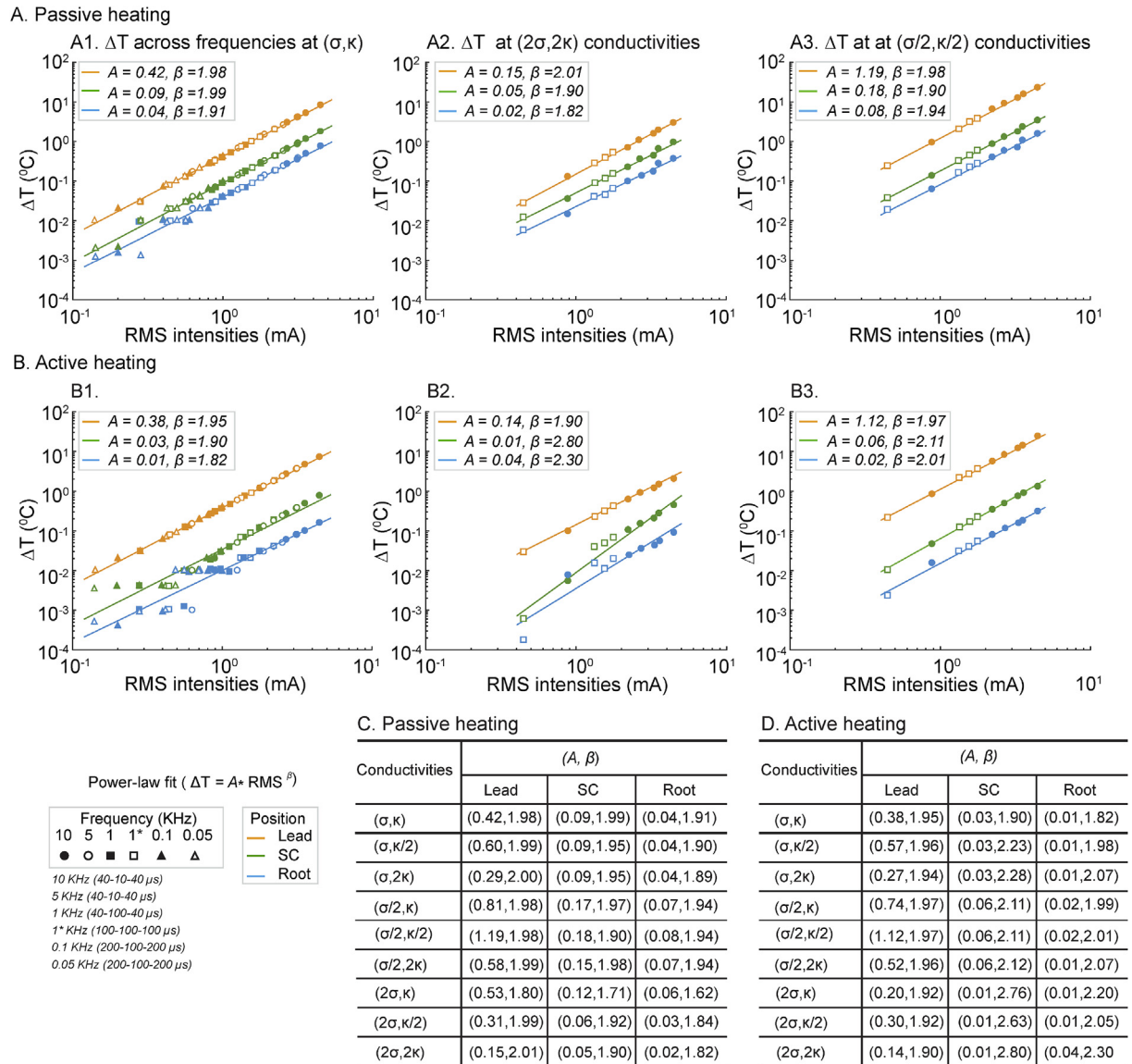


Fig. 4. Power law-fit description of temperature increase with stimulation RMS in SCS model. Temperature changes (ΔT) at three locations (Lead, orange; Spinal Cord, green; and Root, light-blue) as a function of stimulation RMS, across passive or active tissue model with varied fat electrical and thermal conductivity, were predicted by the FEM SCS model (data points). For each tissue model and compartment, we determined a best power-law fit (see Methods), which is a line with proportionality constant A and slope β (lines) on a log-RMS versus log-Temperature plot. Data plots include standard passive (A1), standard active (B1) tissue models, high thermal and electrical fat conductivity ($2\sigma, 2\kappa$) passive (A2) and active (B2) models, and low thermal and electrical fat conductivity ($\sigma/2, \kappa/2$) passive (A3) and active (B3). A and β across all 9 permutations of fat electrical (σ) and thermal (κ) conductivity are summarized for passive (C) and active (D) cases. RMS values plotted in A1 and A2 span stimulation frequencies of 0.05, 0.1, 1 (with low or high duty cycle), 5, and 10 KHz (waveforms indicated in legend and as used in Table 1). RMS values in A2, B2, A3 and B3 span 1, 5, and 10 KHz. $\beta > 1$ indicates super-linear dependence of temperature rise on stimulation RMS; when $\beta = 2$ temperature increases with the square of RMS. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

Cord) and exterior (Lead, Root) regions - indicating that, compared to fat, the Meninges conduct heat away. The reduction in electric field at the Spinal Cord following addition of Meninges (from 165 V/m to 29.27 V/m) was comparable in scale to the temperature decreases in the active model (from 1.22 °C to 0.25 °C) while in the passive model temperature was less sensitive (from 1.37 °C to 0.92 °C) - reflecting that the Meninges are vascularized in the active model. Further addition of CSF (CS) decreased predicted temperature rises at the SC and Root for the passive model, increased predicted temperature rises at the Lead for the passive model, and increased temperature in all compartments in the active model. The avascular nature of the CSF layer is overshadowed by its high electrical/thermal conductivity. Finally, addition of Spinal Cord (SC) restores the tissue parameters of standard model.

Discussion

Thermoregulation of CNS temperature is complex and depends on a high metabolic activity [46] and both passive (conduction) and active heat exchange (blood flow). Neurostimulation, such as SCS, can challenge this equilibrium in several ways by 1) altering neuronal and so metabolic activity [18,47,48]; with 2) changing the cellular microenvironment [48,49]; 3) changing vascular function as a result of both direct blood vessel stimulation [24,50,51] and secondary to microenvironment changes; and 4) depositing of joule heat [18,28]. In the context of kHz-SCS, this study specifically addressed joule heat with the hypothesis that by increased power (pulse compression), kHz-SCS waveforms will superlinearly increase tissue temperature, potentially inducing downstream

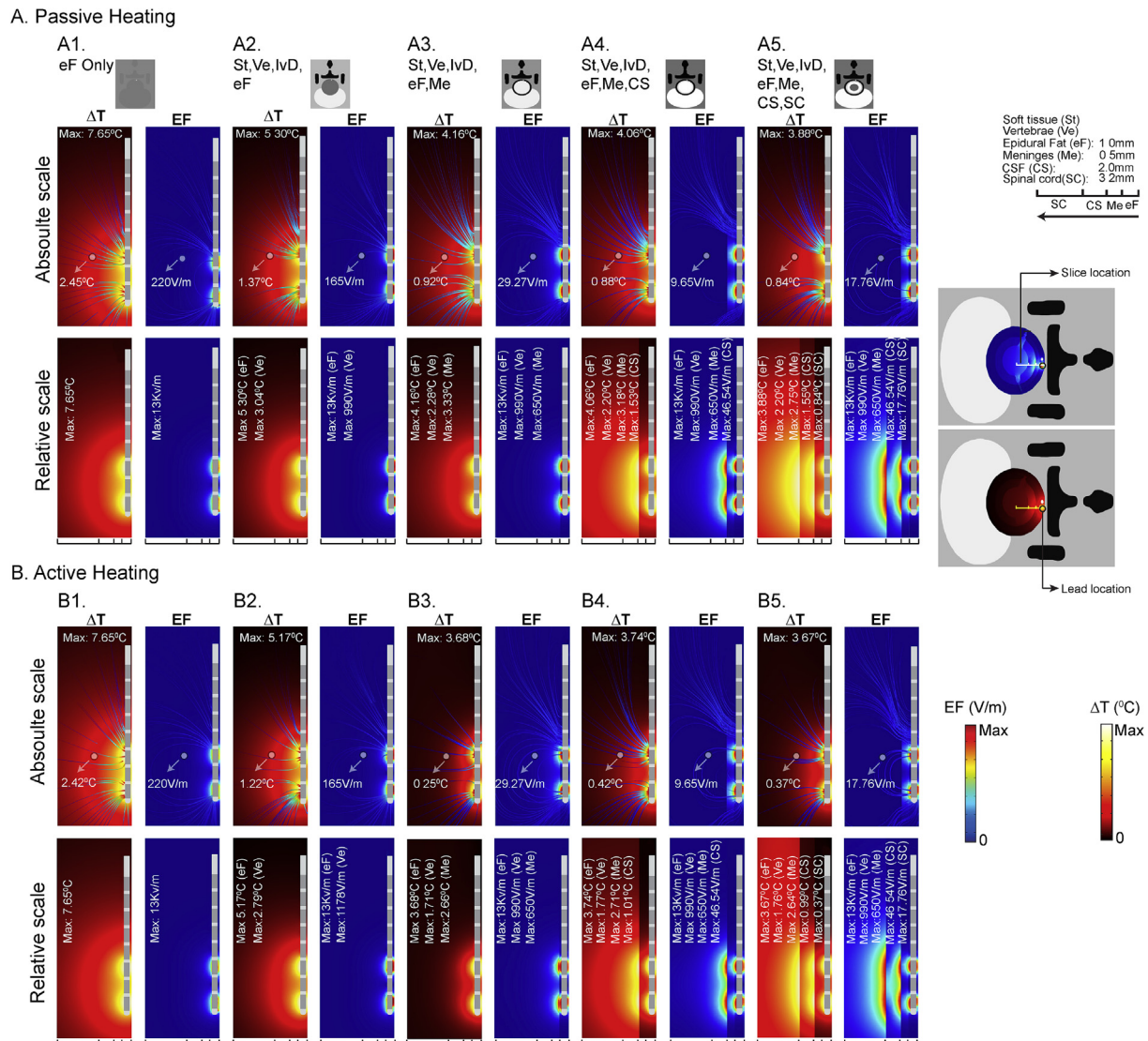


Fig. 5. Tissue substitution analysis for passive and active heating conditions in SCS model. Predicted ΔT and EF were reported in both absolute scale (data sampled at a radial distance equal to the surface of spinal cord from the surface of epidural fat, represented by a gray dot) and relative scale (data sampled at a corresponding tissue surface for all tissue combination as illustrated in the figure) from parasagittal spinal tissue slice. On an absolute scale for both passive and active heating conditions, ΔT and EF decreased as more tissues were added around the epidural fat. With an entire tissue combination (A5, B5) on this scale, the ΔT and EF were 0.84 °C and 17.76 V/m in passive heating and 0.37 °C and 17.76 V/m in active heating. Similarly, on the relative scale for passive and active heating conditions, ΔT and EF decreased as more tissues were added. Maximum ΔT (7.65 °C) and EF (9.9 kV/m) were reported around the lead with only epidural fat (A1, B1, relative scale). As more tissues were added, ΔT around the lead decreased to 3.88 °C in passive heating and 3.67 °C in active heating. Maximum predicted ΔT and EF at the spinal cord was 0.37 °C and 17.76 V/m with the entire tissue combination (A5, B5, relative scale).

Table 2

Tissue substitution order and resulting predicted temperature changes (ΔT) at the Lead, Spinal Cord, or Root tissue compartments. In this series only one waveform with 3.14 mA RMS was simulated. For each model the tissues modeled could include epidural fat (eF), vertebrae (Ve), intervertebral discs (IvD), meninges (Me), cerebral spinal fluid (CS), and spinal cord (SC).

	Passive Heating			Active Heating		
	ΔT (°C)			ΔT (°C)		
Tissues	Lead	SC	Root	Lead	SC	Root
eF	7.39	2.45	0.99	7.32	2.42	1.00
eF, Ve, IvD	5.07	1.37	0.49	5.02	1.22	0.38
eF, Ve, IvD, Me	4.09	0.92	0.37	3.49	0.25	0.03
eF, Ve, IvD, Me, CS	4.13	0.88	0.36	3.59	0.42	0.13
eF, Ve, IvD, Me, CS, SC	4.11	0.84	0.35	3.60	0.37	0.08

alterations in tissue function with therapeutic effects in chronic pain. Characteristic clinical responses to kHz-SCS including as the

lack of associated neural sequelae such as paresthesia and the frequency insensitivity of efficacy [52] reconcile well with joule heating, while the delayed time course of effects [53] may be explain by temperature homeostatic responses or heat shock protein regulation of neuroinflammation.

Bioheat SCS model

FEM bioheat models of phantom, verified experimentally, and of human spinal cord, subjected to a broad parametric sweep (>100 simulations in this study), are suitable for assessing our hypothesis as they enable predictions as to whether or not SCS may produce temperature rises sufficient to produce biological effects.

Heating from chronic SCS represents an exogenous non-physiological challenge. We predicted temperature increases at the dorsal spinal cord of 0.18–1.72 °C and at the lead in epidural fat

of 1.25–11.57 °C under a typical kHz-SCS setting (10 KHz, pulse at 3.5 mA peak; corresponding to 3.13 mA RMS; Fig. 3, Table 1). This range depends on epidural fat electrical conductivity; the combination of high current density and low conductivity increases joule heating that is then conducted to other tissues.

The degree of heating is a super-linear function of stimulation RMS power (Fig. 1D2) such that kHz-SCS can produce significantly more temperature rise than conventional frequency SCS. Assuming $\beta \sim 2$ and integrating (7) with the power-law relationship (4) yields:

$$\Delta T = 0.02 * A * I_{peak}^2 * PCF^2 \quad (8)$$

where '0.02' is the square of RMS_{50} at 1 mA.

Remarkably, at least across conditions considered here, temperature increases in any tissue inside the spinal canal were well fit using a power-law function (equation (8)). With all lead position, electrode configuration, and passive and active tissue properties captured by a single proportionality constant (A ; Fig. 4). All waveform parameters collapse to Pulse Compression Factor, PCF (Equation (7)). This heuristic finding has important potential consequences to SCS practice:

- i. From a modeling standpoint, this finding could dramatically simplify future efforts to predict temperature changes as part of SCS therapy optimization and programming.
- ii. Moreover, the super-linear sensitivity to PCF warrant attentions as incremental changes in waveform can spike tissue heating.
- iii. From a mechanistic standpoint, if temperature increases underpin kHz-SCS, then waveform power (as captured by PCF) is more important than any single waveform parameter (e.g. frequency, pulse width, shape) in generating effective therapy.
- iv. However, a heating MoA does not indicate only waveform PCF predicts outcomes as other factors (e.g. electrode placement) influence the proportionality constant (A). Nor does this imply a fixed minimum for stimulation energy (charge, battery consumption) which depends on other factors such as device efficiency and impedance.

Physiological implications

The nervous system, including the spinal cord, is sensitive to temperature changes. Temperature increases to ~44 °C [51] result in brain damage in animal models after 60 min [36,54], with the temperature threshold for injury decreasing with increased exposure time. In animal models, significant changes in brain excitability have been noted with short-term increases of >2 °C [55–57], with sensitivity to lower-temperature expected with long-term temperature increases. Brain temperature increases above 39 °C in ischemic brain injuries increases extracellular excitatory amino acids level, opening of blood-brain barrier, and elevated proteolysis of the neuronal cytoskeleton [46,58]. A sustained 1–2 °C rise in brain temperature after injury is potentially hazardous [58–60]. While there are *transient* changes in temperature during normal function (2–3 °C [46,58,60,61]) a *sustained* temperature change may produce cumulative and profound changes in brain function. We predicted significant temperature changes in the spinal cord that met or exceeded these thresholds, specifically using kHz frequency waveforms where Pulse Compression increases heat deposition. Our findings are a surprising and important first step toward determining a new heating mechanism for kHz-SCS as well as other relatively high power (kHz frequency) neuromodulation techniques [62,63].

Evidence for stimulation acutely changing neuronal firing and metabolism, perfusion [18,46,47] and the extracellular environment [48] is specific (limited) to sub-kHz frequencies for SCS-relevant stimulation amplitudes [24,49,50]; and so were not modeled here (Q_{met} and w_b were constant). Starting with kHz-stimulation joule heating, changes in brain function can derive from the acute changes in dynamics (e.g. ion channel gating, neurotransmitter clearance; [64,65]) or a homeostatic molecular response to chronic temperature changes (e.g. heat shock proteins). Slow temperature homeostatic changes provide a plausible explanation for the delayed onset of pain relief by kHz-SCS [52,53] and suggest specific molecular pathways (MoA) for pain relief including heat shock protein producing downregulation of neuro-inflammation. For example, 72-kDa heat shock protein (Hsp70) inhibits activation of the pro-neuroinflammatory transcription factor, nuclear factor- κ B in satellite glial cells (NF- κ B) [66]. Knocking out NF- κ B dependent satellite glial cell activation reduces expression of neuronal colony stimulating factor 1 (Csf1) [67], which can potentially reduce the inflammatory response and restore normal function of the spinal pain processing network. However, experimental measurement of expression factors associated such metabolic, genetic, and plastic changes that evolve with prolonged heating and the correlation of these changes with the function of the spinal pain processing network are required to validate this hypothesis.

Conflicts of interest

The City University of New York (CUNY) has IP on neuro-stimulation system and methods with author, Niranjana Khadka and Marom Bikson as inventors. Dr. Marom Bikson has equity in Soterix Medical Inc. Tianhe Zhang, Rosana Esteller, and Brad Hershey are employees of Boston Scientific Inc.

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