Paresthesia Thresholds in Spinal Cord Stimulation: A Comparison of Theoretical Results with Clinical Data

Johannes J. Struijk, Jan Holsheimer, Giancarlo Barolat, Jiping He, and Herman B. K. Boom

Abstract—The potential distributions produced in the spinal cord and surrounding tissues by dorsal epidural stimulation at the midcervical, midthoracic, and low thoracic levels were calculated with the use of a volume conductor model. Stimulus thresholds of myelinated dorsal column fibers and dorsal root fibers were calculated at each level in models in which the thickness of the dorsal csf layer was varied. Calculated stimulus thresholds were compared with paresthesia thresholds obtained from measurements at the corresponding spinal levels in patients. The influences of the csf layer thickness, the contact separation in bipolar stimulation and the laterality of the electrodes on the calculated thresholds were in general agreement with the clinical data.

I. INTRODUCTION

In order to understand the immediate effects of epidural spinal cord stimulation (ESCS), the potential field produced by electrical stimulation and the response of neural elements in the spinal cord to the imposed field should be known. Therefore, modeling may be helpful to gain insight into the mechanisms underlying the phenomena observed in patients, to aid the development of new stimulation methods (e.g., the design of electrodes) and to provide guidelines for further research.

Only a few ESCS modeling studies have appeared in the literature. 2-D calculations of the potential field were carried out by Rusinko et al. [1], Coburn [2] and Sin and Coburn [3]. These calculations may be useful to investigate which geometrical parameters and tissue properties are the most critical. However, when the fields are applied to nerve fiber models 3-D solutions are indispensable.

Three-dimensional calculations were performed by Coburn and Sin [4] and Struijk et al. [5], whereas Swiontek et al. [6] did some field measurements in the spinal cord (post mortem). A major drawback in the latter work is that the stimulating current was applied using electrodes at the pial surface, thus bypassing the shunting cerebrospinal fluid (csf) which, according to the theoretical work [2]-[5], plays a major role in the current distribution.

Coburn [7] applied a calculated 3-D field to a cable model [8] of myelinated dorsal column (DC) fibers and myelinated dorsal root (DR) fibers. He found that the stimulus thresholds for these fibers were of the same order of magnitude as paresthesia thresholds observed in patients and that DR fibers had relatively low thresholds in comparison to DC fibers. This was partly confirmed by Struijk et al. [9]-[10] who calculated thresholds using a more realistic DC fiber model and various DR-fiber models as well.

The relatively low DR fiber thresholds could account for the occurrence of segmental effects, such as a band of tightness around the chest in midthoracic stimulation [11], presumably due to DR fiber activation. These modeling studies also predict that in case of a thin csf layer between electrode and spinal cord, thresholds of DC fibers and DR fibers are approximately the same, which might explain the observations that in some cases paresthesia was first perceived in the lower extremities [11] during cervical stimulation.

In the present study, models of ESCS, comprising volume conductor models of the spinal cord and surrounding tissues at midcervical, midthoracic and low thoracic vertebral levels, and models of DC fibers and DR fibers were used to simulate clinical data. The impedance between bipolar electrodes in the volume conductor models was adapted to the load impedances measured in patients. In this initial validation study the influences of the dorsal csf layer thickness, the contact separation in bipolar stimulation and the laterality of the electrodes on stimulus threshold were investigated. The calculated stimulus thresholds were compared with paresthesia thresholds measured in patients.

II. METHODS

A. Volume Conductor Models

Three-dimensional volume conductor models of the spinal cord at midcervical, midthoracic, and low thoracic vertebral levels were used in this study. Transverse sections of these models are shown in Fig. 1. Each model comprises the spinal cord, which is composed of gray matter (gm) and white matter (wm), cerebrospinal fluid (csf), epidural space (es), vertebral bone (vb), a layer representing surrounding tissues (sl), electrode contact insulation (is) and a thin layer representing the dura mater (dm) at the dorsal side. The small dorsal root filaments, immersed in the well-conducting csf, were not incorporated in the volume conductor model [10].

The tissue conductivities used in the model are given in Table I. The values of the white matter, gray matter, epidural fat and vertebral bone were taken from Geddes and Baker [12]. We measured the csf conductivity at 37°C previously...
in samples from three subjects [5]. The conductivity of the dura mater is unknown. Its model value was taken such that the total impedance between two contacts (10 mm center separation) matched the impedance measured bipolarly in patients (mean = 109Ω, sd = 346Ω, n = 114). Therefore, the dura impedance also includes the impedance of the tissue (connective tissue, epidural fat) which may be present between electrode and dura. In simulations of unipolar (cathodal) stimulation the boundary of the model serves as the distant contact. The conductivity of the surrounding layer (sl) was given a value such that the impedance between cathode and boundary matched the impedance measured unipolarly (mean = 715Ω, sd = 229Ω, n = 69). In the model the impedance was obtained by calculating the currents at the surface of each contact.

The contacts were modeled as voltage sources (Dirichlet condition) because commercially available stimulators have voltage sources. The dimensions of the contacts were 3.6 × 3.6 mm, matching the contact areas of the Resume® lead (Medtronic, Inc., Minneapolis, MN). Bipolar configurations with center separations of 10, 20, 30, 40, and 50 mm and a unipolar configuration were used. Because of the different morphology distinct vertebral levels had to be modeled separately. In relation to the availability of clinical data the following vertebral levels were chosen: midthoracic (C4–C6), midthoracic (T4–T7) and low thoracic (T10–T11).

The dorsal csf layer thickness at each level was estimated from an MRI study on healthy subjects (18 males, 20–40 years of age). The values of the median and the 25th and 75th percentiles are given in Table II. The 95% confidence intervals were calculated according to the method given in the Appendix, which is a modification of the method used to calculate the confidence interval of the median [13].

In Fig. 1 for each level a model is shown with the median value of the dorsal csf layer thickness at that level. Similar models were used with csf layer thicknesses equal to the 25th percentile and the 75th percentile of the MRI data (Table II). These models were used for simulating the clinical data. Because paresthesia thresholds had a non-Gaussian, skew distribution, the use of the median and the percentiles was preferred to the use of the mean and standard deviation to characterize the distributions.

The dimensions of the spinal cord are given in Fig. 1 and were taken from literature [14]–[20]. For the low thoracic level only a few data were available. The cord morphology at all three vertebral levels was taken from Fix [21]. The length of the model was 60 mm.

To discretize the volume conductor model, a rectangular grid was used [see Fig. 1(a)]. Grid spacings varied from 0.2 to 1.6 mm with the smallest values near the electrodes and the dorsal columns. The number of grid points was 185193 (57 × 57 × 57). A finite difference method (using Taylor series) was applied to discretize the governing Laplace equation. The resulting set of linear equations were solved using a Red–Black Gauss-Seidel iteration [22] with variable overrelaxation (overrelaxation after each red and each black sweep). Two solutions were calculated simultaneously, one starting with an underestimated initial solution and the other one with an overestimated initial solution. The overrelaxation factor was increased if the two solutions approached each other too slowly, whereas it was decreased if the difference between the solutions decreased too quickly, thereby avoiding
The calculations were finished when the average absolute difference between the two solutions was less than 0.01% of the average absolute value of the calculated field itself. For a special case including discontinuous conductivities, which could also be solved analytically, the numerical solutions appeared to be accurate within 2% [10].

B. Nerve Fiber Models

Two types of nerve fiber models were used: DR fibers and DC fibers. DC fibers are longitudinal fibers in the dorsal columns of the gray matter (adapted from Cajal [23]); (b) Network model consisting of a rostrocaudal fiber (horizontal part) and a collateral (vertical part); $R_m$: nodal membrane resistance, $C_m$: nodal membrane capacitance, $R_e$: internodal intracellular resistance, $V$: nodal intracellular potential, $V_i$: extracellular (applied) potential.

instability of the iteration process due to the overrelaxation. The calculations were finished when the average absolute difference between the two solutions was less than 0.01% of the average absolute value of the calculated field itself. For a special case including discontinuous conductivities, which could also be solved analytically, the numerical solutions appeared to be accurate within 2% [10].

B. Nerve Fiber Models

Two types of nerve fiber models were used: DR fibers and DC fibers. DC fibers are longitudinal fibers in the dorsal columns issuing collaterals into the dorsal gray matter (adapted from Cajal [23]). The model of these fibers is a cable model [8] in which the myelin is assumed to be a perfect insulator, extended with collaterals as shown in Fig. 2(b). All nodes were madeexcitable using the equations by Chiu et al. [26]transformed to a temperature of 37°C. The collaterals were attached to every second node of Ranvier of the main fiber. This model was described in detail in a previous paper and simulations indicated that the presence of collaterals reduces the stimulus threshold (the minimum stimulus amplitude at which a given fiber is excited) by 30–50% [9]. A stimulus pulse width of 210 μs was used to calculate the stimulus thresholds of both DC and DR fibers.

For the DR fiber we used a cable model with a curved trajectory, the proximal end being connected to a DC fiber model. The curvature of the model is shown in Fig. 3. The model and its properties were described previously [10]. The fiber entered the spinal cord at a rostrocaudal level corresponding to the center of the cathode, at which position the stimulus threshold had the lowest value.

The fiber models were given diameters in the upper range of the diameter distribution of these fibers. We assumed that activation of the hair receptor fibers (Aαβ fibers [25]) is responsible for paresthesia perception, because they are the largest fibers and will thus have the lowest excitation thresholds. We used a DR fiber diameter of 15 μm. Assuming a linear relationship between propagation velocity and fiber diameter [27], the attached DC fiber was given a diameter of 12 μm, in accordance with Desmedt and Cheron [28] who found a drop in propagation velocity of about 19% after bifurcation of the DR fiber in the human dorsal columns.

The transverse position of the DR fiber in the volume conductor model is shown in Fig. 1(c). The positions of two DC fibers, indicated by 1, 2, are also shown. Fiber 1 has a dorso-medial position and is therefore assumed to be a fiber of sacral origin [24], [29], [30]. In accordance with Petit and Burgess [31], who found a decrease in propagation velocity of 40–65% along the ascending fibers in the dorsal columns of the cat, fiber 1 was given a diameter of 7.5 μm. This value corresponds to the upper range of fiber diameters in the human fasciculus gracilis as found by Ohnishi et al. [32] (up to 8 μm), although Häggqvist [33] found fiber diameters of more than 10 μm in this area. Fiber 2 represents a DC fiber close to the entry level of the corresponding DR fiber and had a 12 μm diameter. We will refer to these fibers as fiber DC1, DC2 and DR.

C. Collection of Clinical Data

The impedance values and paresthesia thresholds in ESCS were measured in patients with either an Itrel® or Itrel II® implanted pulse generator with a Resume® lead (Medtronic Inc., Minneapolis, MN). The 108 patients included in this study were suffering from either chronic pain or spasticity. The main aetiologies were reflex sympathetic dystrophy, low back pain and spinal cord injury. Paresthesia threshold was defined as the minimum voltage at which the patient perceived paresthesia (a tingling or buzzing sensation due to activation of afferent hair fibers), measured with the patient in the supine position at a repetition rate of 50 pps and a pulse width of 210 μs.

The stimulation amplitude was gradually increased starting from 0 V with increments of 0.25 V (Itrel®) or 0.1 V (Itrel II®) to detect paresthesia threshold. At each increment the patient was asked to report whether and where paresthesiae were perceived. Impedance was measured at 50 pps, a pulse width of 210 μs and an amplitude of 1.0 V using the impedance measurement option of the Itrel II® implantable pulse generator.

The Resume® lead is an array of four circular contacts embedded in a strip of flexible insulating material. The contacts have a diameter of 4.0 mm and a center separation of
Measurements were carried out with a unipolar electrode configuration (the metal case of the pulse generator being the anode) and bipolar configurations with 10, 20, and 30 mm contact separations as well.

Data were obtained from patients having the electrodes placed between 0 and 3 mm lateral to radiological midline at the vertebral levels C4–C6 (31 patients), T4–T7 (24 patients) and T10–T11 (53 patients). The radiological midline is related to the spinous processes and does therefore not necessarily match the vertebral levels C4–6 (31 patients), T4–T7 (24 patients) distributed the median and the 25th and 75th percentile were calculated. The 95% confidence intervals were established using the method given in the Appendix.

Because the paresthesia threshold data were not Gaussian distributed the median csf layer thickness the measured paresthesia thresholds are mainly due to differences in dorsal csf layer thickness. The model we used csf layer thickness values equal to the measured thresholds (see Discussion). We calculated the effect of csf layer thickness on threshold and compared the variation in calculated thresholds due to the variation in csf layer thickness with the observed variation in paresthesia thresholds, measured with the patients in supine position. Because the dorsal csf thicknesses of the patients were not measured and consequently unknown, this comparison makes sense if it is assumed that differences in paresthesia thresholds are mainly due to differences in dorsal csf layer. We indeed made this assumption on the basis of the results on the relationship between paresthesia threshold and spinal level, and the high sensitivity of thresholds to a patient’s position.

It is well known that paresthesia threshold varies with a patient’s position. A supine position yields lower thresholds than an upright position. In cervical stimulation the set of the head strongly influences threshold. Both the patient’s position and the set of the head affects the dorsal csf layer thickness. Because the paresthesia threshold data were not Gaussian distributed the median the 25th and 75th percentile were calculated. The 95% confidence intervals were established using the method given in the Appendix.

III. RESULTS

A. Paresthesia Threshold as a Function of Spinal Level

The dorsal csf layer thickness is a major parameter affecting the threshold stimulus for the excitation of nerve fibers in the spinal cord, because this layer determines the electrode-fiber distance [5], [10], [34]. Table II shows that at the midcervical level the dorsal csf layer thickness is smallest and therefore it is expected that paresthesia thresholds will have the lowest values at that level. We calculated the stimulus threshold $V_{th}$ of the fibers DC1, DC2 and DR at the three levels in bipolar stimulation with a contact separation of 10 mm and a dorsal csf layer thickness being the median of the measured values (Table II). The minimum values of $V_{th}$, being $V_{th}$, of the DR filter in all cases, are summarized in Table II. In the midcervical model with median csf layer thickness the values of the fibers DC1 and DC2 were about 100 and 50% higher, respectively. At midthoracic and low thoracic levels these differences were even more pronounced. Therefore, the expected recruitment order is DR fiber, lateral DC fiber and medial DC fiber successively.

For very small csf layer thicknesses (less than 1.0 mm) this order was reversed. In Table II the midcervical model predicts that for DR fibers a minimum $V_{th}$ occurs at a contact separation of about 20 mm. Simulations with 30, 40, and 50 mm contact separation showed a continuous slight increase of $V_{th}$ (11% increase from 30 mm to 50 mm). For DC fibers however, minimum $V_{th}$ occurred below 10 mm.

B. Paresthesia Threshold as a Function of csf-Layer Thickness

We calculated the effect of csf layer thickness on threshold and compared the variation in calculated thresholds due to the variation in csf layer thickness with the observed variation in paresthesia thresholds, measured with the patients in supine position. Because the dorsal csf thicknesses of the patients were not measured and consequently unknown, this comparison only makes sense if it is assumed that differences in paresthesia thresholds are mainly due to differences in dorsal csf layer. We indeed made this assumption on the basis of the results on the relationship between paresthesia threshold and spinal level, and the high sensitivity of thresholds to a patient’s position.

In Fig. 4 the median and the 25th and 75th percentiles of the measured paresthesia thresholds $V_{th}$ normalized to the median values at that level. We calculated the stimulus threshold $V_{th}$ of the fibers DC1, DC2, and DR using bipolar stimulation with a contact separation of 10 mm. The results for the DR fiber, normalized to the values with median csf layer thickness, are shown in Fig. 4 (bars). In all cases fiber DR had the lowest $V_{th}$ value, although in the midcervical model with a 25th percentile csf layer thickness fibers DC1 and DC2 had only slightly higher values (40 and 20%, respectively).

C. Paresthesia Threshold as a Function of Contact Separation

We calculated the stimulus thresholds $V_{th}$ of fibers DC1, DC2, and DR in the three volume conductor models with median csf-layer thickness and contact separations of 10–50 mm, with steps of 10 mm and a unipolar configuration as well. The DR fiber again had the lowest $V_{th}$ value. These values (except for the 40 and 50 mm separation), normalized to $V_{th}$ at 10 mm contact separation are summarized in Fig. 5 (bars). These data show that in bipolar stimulation the model predicts that for DR fibers a minimum $V_{th}$ occurs at a contact separation of about 20 mm. Simulations with 30, 40, and 50 mm contact separation showed a continuous slight increase of $V_{th}$ (11% increase from 30 mm to 50 mm). For DC fibers however, minimum $V_{th}$ occurred below 10 mm.
contact separation as we have shown previously [34], [35]. For fiber DC1 at the midcervical level normalized $V_{th}$ values were 100% (3.37 V), 112%, and 117% for 10, 20, and 30 mm contact separation, respectively, whereas for fiber DC2 these values were 100% (2.56 V), 114% and 119%. So, in our model, we found that with increasing contact separation (up to 50 mm), the ratio $V_{th}(DR)/V_{th}(DC)$ decreases, thus increasing the preferential stimulation of DR fibers. The highest preference to DR fibers was obtained in unipolar stimulation.

Barolat et al. [11] found that overall paresthesia threshold increased slightly as the contact separation was increased from less than 30 mm to more than 50 mm, whereas the percentage of body area covered by paresthesia decreased with increasing contact separation. This is well in accordance with our theoretical results if we assume that in dorsal root stimulation a smaller percentage of body area will be covered by paresthesia than in dorsal column stimulation, in which case fibers originating from lower spinal levels are activated as well.

Because of the relatively large confidence intervals the measurement data as shown in Fig. 5 (markers) do not show a significant change (Kruskal–Wallis test [13], $p = 0.05$) of paresthesia threshold as contact separation is increased from 10 mm to 30 mm (see [11] for larger distances). The slight minimum which occurred at 20 mm in the calculated $V_{th}$ was not observed in the experimental data.

In the model, unipolar stimulation resulted in a relatively low $V_{th}$. The measurement data for unipolar stimulation show large confidence intervals due to the small number of patients in which unipolar measurements were performed (10, 5, and 17 patients at the midcervical, midthoracic and low thoracic level, respectively).

**D. Influence of the Lateral Position of the Electrode Contacts**

According to Barolat et al. [11] thresholds and paresthesia distributions are largely affected by the laterality of the contacts. They measured an average paresthesia threshold of 1.7 V for all bipolar combinations with contacts within 3 mm of the radiological midline at thoracic vertebral levels. The average threshold dropped to 1.2 V (70%) for contacts at 3–5 mm from the midline and 0.8 V (47%) for contacts at more than 5 mm from midline. This effect is most likely related to dorsal root stimulation. With both contacts at one side at 0.5 to 3 mm from midline (see Fig. 6), most paresthesia distributions were unilateral. When the anode was more than 0.5 mm from midline contralateral to the cathode, bilateral paresthesia distributions slightly increased. However, the position of the cathode was the most critical [11].
Fig. 6. Positions of the contacts in a coronal plane. (a) Contacts at midline; (b) contacts at the same side; (c) cathode lateral, anode at midline; (d) contacts at opposite sides.

<table>
<thead>
<tr>
<th>CONTACTS</th>
<th>DC1</th>
<th>DC2-left</th>
<th>DC2-right</th>
<th>DR-left</th>
<th>DR-right</th>
</tr>
</thead>
<tbody>
<tr>
<td>Midline</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Same side</td>
<td></td>
<td>(13.9 V)</td>
<td>(10.0 V)</td>
<td>(10.0 V)</td>
<td>(4.20 V)</td>
</tr>
<tr>
<td>Cathode</td>
<td>1.44</td>
<td>1.41</td>
<td>1.64</td>
<td>0.90</td>
<td>1.76</td>
</tr>
<tr>
<td>Opposite</td>
<td>1.40</td>
<td>1.60</td>
<td>0.94</td>
<td>1.68</td>
<td>0.85</td>
</tr>
</tbody>
</table>

In order to assess the relative influence of the anodal and cathodal position we did four simulations (see Fig. 6): (1) both contacts at midline, (2) both contacts at 3.0 mm from midline, (3) cathode at 3.0 mm from midline at the right side and anode at midline, (4) cathode at 3.0 mm at the right side and anode at 3.0 mm left to midline. The mid-thoracic model was used with medial dorsal csf layer thickness of 1.75 mm, with thresholds measured at midthoracic and high thoracic levels [36], where in reality the median of the csf layer thickness is about three times larger. We calculated, that the difference between thresholds for a 1.75 mm csf layer and a 5 mm csf layer is more than a factor 3. Therefore, Cobum’s results show the same discrepancy between modeling and clinical data as ours.

The discrepancy between calculated and measured thresholds may be due to variability of parameters in both the nerve fiber models and the volume conductor models. The geometrical and electrical parameters of mammalian myelinated nerve fibers are described in literature [8], [27], [37]. In our fiber models average parameter values were used [9]. However, most parameters have large standard deviations and therefore, the threshold stimulus of fibers with a given diameter and a given position may vary largely. Wide physiological ranges have been observed for, amongst others, the ratio of the internodal length and the fiber diameter (L/D), ratio of axon diameter and outer fiber diameter, nodal membrane capacitance, nodal membrane conductivity, intra axonal conductivity and length of the node of Ranvier [27], [37]. We calculated (unpublished results) a threshold stimulus distribution of a fiber model where those parameters were given a normal distribution with standard deviations of 20% of their average values (from data in the literature we calculated a standard deviation being 26% of the average value of L/D). We found that about 5% of the distribution was below half of the median threshold value of the distribution and it may be assumed that this 5% (or even less) of “low threshold” fibers defines the threshold of the whole fiber population (of given diameter) in the dorsal columns or dorsal roots. The parameter variability will thus significantly decrease the threshold of a population of nerve fibers in the spinal cord or in the dorsal roots.

The variability of the membrane capacitance hardly had any influence, because of the relatively long pulse (210 μs). A parameter of the volume conductor model which strongly affects stimulus thresholds, is the dorsal csf layer thickness [5]. The number of MRI scan data used to estimate this parameter was relatively small, thus giving rise to the possibility of large errors in the estimates (large confidence intervals, see Table II). The insertion of a relatively large electrode lead in the narrow epidural space will indent the dura, thus decreasing the csf layer thickness and decreasing paresthesia threshold. Another geometrical factor giving rise to a relatively low measured paresthesia threshold is the fact that the electrode position was up to 3 mm off midline.

In this investigation the calculated threshold voltages for the activation of DC and DR fibers of the spinal cord with epidural electrical stimulation were compared with measured paresthesia thresholds in patients. We assumed that these myelinated primary afferent fibers are involved in ESCS and that paresthesiae, at threshold levels, are directly related to activation of these fibers without modulation by excitatory or inhibitory spinal or long loop circuits. At stimulus amplitudes beyond threshold, relations may be more complex.

We found a discrepancy between paresthesia thresholds and calculated stimulus thresholds, the latter being 252–300% of the measured values (Table III), whereas Cobum seemed to have obtained a better fit between the modeling results and measured paresthesia thresholds [7]. However, he compared thresholds obtained with a model with a dorsal csf thickness of 1.75 mm, with thresholds measured at midthoracic and high thoracic levels [36], where in reality the median of the csf layer thickness is about three times larger. We calculated, that the difference between thresholds for a 1.75 mm csf layer and a 5 mm csf layer is more than a factor 3. Therefore, Cobum’s results show the same discrepancy between modeling and clinical data as ours.

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The modeling results show that variations in CSF layer thickness can largely account for the variability of the measured paresthesia thresholds among patients. One would expect the impedance to also affect Vp, but no statistically significant relationship was found between measured impedance and Vp. The influence of the impedance variability on Vp might be overshadowed by the variations in CSF layer thickness. In the model, the CSF layer thickness changes only caused less paresthesia to start at a lower level which can be explained by a physiological range, CSF layer thickness and impedance are hardly correlated.

Paresthesia usually starts in the dermatomes corresponding to the spinal level of the cathode [38] and often spreads caudally with increasing stimulus intensity [11]. This behavior is consistent with the model prediction that recruitment starts in the dorsal roots and spreads into the lateral and the medial dorsal columns with increasing amplitude. Sometimes, paresthesia starts at a lower level which can be explained by a reverse recruitment order as obtained in the midcervical model with a small dorsal CSF layer thickness.

From this initial validation study it can be concluded that a consistent discrepancy exists between the absolute values of the calculated threshold stimuli and the measured paresthesia thresholds at all three spinal levels. Further improvement of the model is needed to eliminate this discrepancy. Therefore, reliable data on the spread of some model parameters, such as the dorsal CSF layer thickness in patients with implanted electrodes and nerve fiber parameters are needed. On the other hand, the change in calculated threshold values as a function of CSF layer thickness, contact separation and laterality were in general agreement with both the clinical measurements and the predicted spread of paresthesia. The model may therefore be used to predict the effects of various electrode configurations. The influence of contact dimensions, electrode position and contact combination on the stimulus threshold of DC fibers and DR fibers can be assessed with the model. Other neural elements, such as fibers in the dorsolateral funiculus and dorsal horn cells can be incorporated into the model. The method described in this paper may thus be a useful tool for designing new electrodes which will make it possible to activate various neural pathways or structures in a more specific way.

V. APPENDIX

A. Confidence Interval of the 100p th Percentile Using the Sign Test

The confidence interval of the estimated 100pth percentile of a random sample X = X1, X2, ..., Xn of nonparametric data can be obtained using the Sign test (for the 25th percentile, the median and the 75th percentile p = 0.25, p = 0.5, p = 0.75, respectively).

Let Π be the 100pth percentile of a random sample X.

Test the null hypothesis H0 : Π = Π0

against the alternative H1 : Π ≠ Π0

using the Sign test statistic S = number of Xi for which Xi < Πi, i = 1, ..., n. Then S has a binomial distribution [21] because the probability of Xi < Πi equals p under the assumption H0. Thus, P(S = s) = B(s, n, p) = (n \choose s) · p s · (1 − p) n−s, for s = 0, 1, 2, ..., n. H0 is rejected at level α if S ≤ s1 or if S ≥ s2 where s1 and s2 follow from:

P(S ≤ s1) = α/2

P(S ≥ s2) = α/2

The latter can be rewritten as P(S ≤ s2 − 1) = 1 − α/2.

For given n, α and p, the values of s1 and s2 can be obtained from a table of the binomial distribution.

Repeating this test procedure for all possible values of Π0 a 100(1−α)% confidence interval is then the range of values Π0 so that Π is in the acceptance region. If the observations X1, X2, ..., Xn are ordered from smallest to largest, the confidence interval becomes

Xs2 + 1 to Xs2.

REFERENCES


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