Early and preventable changes of peripheral nerve structure and function in insulin-deficient diabetic rats

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Summary Structure and function of peripheral nerve in early insulin deficient streptozotocin diabetic rat were examined. Quantitative studies of myelinated fibres from electron micrographs showed that fibre calibre and slope coefficients of regression lines between axon and myelin areas were reduced and quantitation of unmyelinated fibres revealed a 40% reduction in area fraction of smooth endoplasmic reticulum. The difference in conduction velocity between diabetic rats and control rats increased with time just as does the difference in fibre calibre. Insulin treatment was able to prevent the decrease in conduction velocity as well as the reduction in fibre calibre. It is concluded that the early structural axon abnormality results from the diabetic state and cannot be explained by poor general condition of the animals.

After several years of diabetes some patients develop symptoms of peripheral neuropathy (Lundbæk, 1953). Nerve function is considerably disturbed in most long-term diabetics and nerve structure is severely damaged including segmental demyelination and axonal loss (Thomas and Lascelles, 1966; Reske-Nielsen and Lundbæk, 1968; Chopra et al., 1969).

In early insulin-deficiency juvenile diabetics, motor nerve conduction velocity is slightly reduced (Gregersen, 1967). If the metabolic state is normalised or nearly normalised, as can be done for a few weeks during hospitalisation, conduction improves. When the metabolic state deteriorates by withdrawal of insulin treatment, motor conduction is again slowed (Gregersen, 1968a). The same is true for peripheral nerve function during ischaemia (Gregersen, 1968b). These results indicate that the nerve abnormalities demonstrable by these neurophysiological techniques are caused by the metabolic derangement which characterises the diabetic state.

There is very little information on the structural background of these functional changes in early juvenile diabetes. One reason for this is, of course, that physicians hesitate to do nerve biopsies in recently diagnosed young patients without symptoms of peripheral neuropathy.

In experimental insulin-deficiency diabetes, nerve conduction and peripheral nerve function during ischaemia are known to be changed as in early juvenile diabetes (Eliasson, 1964; Seneviratne and Peiris, 1969). Light microscopic studies in our laboratory have shown changes in the fibre calibre corresponding quantitatively to the functional abnormalities (Jakobsen, 1976a; Jakobsen and Lundbæk, 1976).

This conclusion has been criticised by Thomas and Sharma (1976) and Sharma et al. (1977). They suggested that what we had found is not a nerve change caused by the metabolic abnormality characterising the diabetic state of the animals, but just an expression of general poor health with retardation of growth in length.

We have, therefore, repeated our experiments this time including measurements of skeletal growth. The results obtained show that in our experimental arrangements skeletal length is identical in diabetic and control animals.

Furthermore, we have extended our studies of early structural nerve changes by an electron microscopic quantitation of calibre of myelinated
and unmyelinated fibres, axon-myelin ratios and of axon organelles. In order to find out whether the slowing of nerve conduction was correlated with the difference in fibre calibre, conduction velocity was examined with respect to duration of diabetes. Finally, prevention of the structural and functional nerve abnormalities occurring in streptozotocin diabetic rat was attempted by insulin treatment.

Materials and methods

Male Wistar rats were used throughout these experiments. With the exceptions noted below all the rats were 28 weeks old at the time of sacrifice or determination of nerve conduction velocity.

For the study of age changes in normal rats, groups of animals were examined at the age of 24 and 28 weeks (body weight 350–450 g), and at the age of 48 weeks (body weight 360–505 g).

Skeletal growth in diabetic animals was studied in a group of 10 animals aged 24 weeks which had been diabetic for four weeks. An age- and weight-matched group of 10 control rats was used for comparison (body weight 330–380 g at the age of 20 weeks). Skeletal growth was also studied in the group of 48 week old rats (n=10) mentioned above.

The effect of insulin on nerve structure was studied in a group of rats which had been treated with insulin for four weeks from the induction of diabetes at the age of 24 weeks.

Ultrastructural investigations were performed at electron microscopy on material available from our earlier study (Jakobsen, 1976a). The rats were 23–24 weeks old and had had diabetes for four weeks. Age- and weight-matched control rats were used for comparison (body weight 330–380 g at the age of 19–20 weeks).

Nerve conduction velocity was determined in four groups of diabetic rats, all aged 28 weeks, but with a duration of diabetes between 10–18 hours and five weeks. A control group and a group of diabetic rats treated with insulin for two weeks were also included in these neurophysiological studies (body weight 350–450 g at the time of selection for the experiment).

Diabetes was induced by intravenous injection of 40–45 mg streptozotocin per kg body weight, and only rats having a blood glucose value above 13.9 mmol l⁻¹ (250 mg/dl) 24 hours after the injection entered the experiments. Blood glucose was determined with Dextrostix and an Ames reflectance meter.

A long-acting insulin preparation (modified Ultralente, pH 5.5), kindly provided by Dr J. Schlichtkrull, Novo Research Institute, was used in this study (Rasch, 1979). Insulin was given daily at noon, and the dose was selected according to blood glucose determinations made just before the injection. The first insulin dose was given 24 hours after the application of streptozotocin and the last one less than 24 hours before the final experiments. The insulin dose given within the first week was reduced gradually from 8–4 iu per day till it was stabilised at about 3 iu per day. It was intended to keep the blood glucose as close to normal values as possible. Values between 3.3 and 6.7 mmol l⁻¹ (60 and 120 mg/dl) were accepted without leading to a change of the insulin dose. Blood glucose was determined every two hours in a 24 hour period two weeks after the start of the experiment. Five animals with values between 3.3 and 6.7 mmol l⁻¹ that day were selected for the experiment.

Skeletal growth was estimated by measuring tibial length. The tibia was freed from connective tissues, the cartilages were left intact and the measurement of the wet bone was performed with calipers.

Light microscopy

The common peroneal nerves were fixed by vascular perfusion, through the abdominal aorta under controlled pressure, with a cacodylate buffered paraformaldehyde-glutaraldehyde fixative. The osmolarity of buffer and fixative was measured by freezing point depression and maintained within narrow limits. A 10–20 mm long specimen of the nerve was taken just above the knee, postfixed in 2% osmium tetroxide, rinsed in buffer, dehydrated in graded acetone and embedded in Vestopal. Thin transverse sections (0.1–0.2 μm) were cut on an LKB-Ulrotome and stained with toluidine blue. Transverse sections were obtained by placing the nerve axis perpendicular to the glass knife by means of a 10 times magnifying ocular fitted with cross hairs.

To obtain representative samples of central as well as of peripheral parts of the nerve, cross-section sampling was made within circle sectors placed at random on the image of the section. Myelinated fibres were counted and measured within two sectors making up one-eighth of the nerve area. Fascicular area, endoneurial area, sector area, and fibre area were all estimated by a point-counting technique (see Elias et al., 1971; Weibel and Bolender, 1973) at magnifications and point densities as previously described by Jakobsen (1976a). The endoneurial area in this part of the study means the fascicular area minus the myelinated fibre area and the vessel area (see Jakobsen, 1978).
The reason why area estimation of fibre calibre was performed instead of measuring “fibre diameter” is that the cross-sectional area is an unbiased estimate whereas “fibre diameter” is a rough estimate because fibre profiles are neither circles nor ellipses. An important consequence of this often neglected fact is that measurements of “diameter” may lead to errors when fibres of different shapes are compared. In such situations, the fibre “diameter” may change in a way inconsistent with the relation between diameter and area. This, of course, may well be the case in studies of pathological conditions.

**Electron Microscopy**

The nerves used for the ultrastructural study were fixed, dehydrated, and embedded as described above. A sample corresponding to about 14% of the fascicular area was obtained by trimming a small pyramid which did not include the centre of the nerve. Ultrathin sections of the sample were cut on an LKB-Ultratome and stained with uranyl magnesium acetate and lead citrate. Electron microscopy was carried out with a Philips EM 200, and the micrographs of the selected part of the common peroneal nerve were pasted together to form one composite picture, which included an average of nearly 250 myelinated fibres. The magnification was checked by photographing a carbon grating grid on each film. Its average value was 13,500 times.

When measuring myelinated fibres on electron micrographs a sampling problem is present. Fibres cut by the border of the micrographs cannot be measured. Since small fibres can be measured closer to the border than larger ones, the smaller fibres are over-represented. This problem was solved by drawing a frame at the distance of one large fibre diameter from the inside of the border of the photomontage. In this way fibres cut by the montage were omitted, while fibres divided by the frame were included in proportion to the part of them which lay inside the frame.

The size of the test area was estimated with a point density of one point per 45 μm², and the number of fibres in the common peroneal nerve was obtained by multiplying the number of fibres within the test area by the ratio between the fascicular area determined at light microscopy and the test area.

Fibre, axon, and myelin areas were estimated with a point density of one point per 5 μm². Only points falling upon compact myelin were counted, excluding incisures of Schmidt-Lanterman and areas of myelin splitting.

The number of mitochondrial profiles and smooth endoplasmic reticulum (SER) profiles within all myelinated fibres in the test area, and the total areas of these organelles were estimated using a point density of one point per 0.13 μm². Mitochondria were defined in this study as dark organelles with inner structure and SER as light organelles without inner structure.

The density of neurofilaments was estimated by counting the number of filaments within 10 small quadrants of 0.09 μm² within each of 10 myelinated axons per animal. Randomised selection of 10 numbered fibres was obtained by means of a table of random numbers.

Measurements of unmyelinated nerve fibres were started in the upper left corner of each photomontage and stopped when the counts exceeded 200 fibres (range, 207–257). The numbers of mitochondrial profiles and SER profiles within these fibres were counted, and fibre size, mitochondrial, and SER areas were estimated with a point density of one point per 0.11 μm².

**Conduction Velocity**

To minimise the variation of length measurements and temperature in determinations of nerve conduction velocity of the largest fibres the tail nerve was used (Miyoshi and Goto, 1973). The rats were anaesthetised mildly with a sodium pentobarbitone-diazepam mixture, and the left ventral tail nerve was exposed surgically 60 and 120–140 mm from the anus, and cut distally.

The tail was lead into a vessel in the wall of which thermostated water was circulating. Subsequently the vessel was filled with paraffin, and the temperature of the paraffin bath was maintained between 37.5°C and 38.5°C. Before the experiments the tail was kept in the bath for 20 minutes reducing the temperature difference between the paraffin and the subcutis of the tail to less than 0.3°C. Then the nerve was placed on J-shaped 0.15 mm thin platinum electrodes and stimulated proximally with single rectangular current pulses. The propagated action potentials were recorded distally. Stimuli with a duration of 0.05 msec supramaximal to the fastest conducting fibres were delivered from a stimulator at a frequency of 10/s. Thirty consecutive action potentials were displayed on a Medelec® oscilloscope, superimposed and photographed with a time scale. The distance between the two pairs of electrodes was measured before and after stimulation with a pair of pointed-leg calipers using the mean value for calculation of conduction velocity.

The rats used in the experiments for ultrashort duration of diabetes had had blood glucose values above 11.1 mmol l⁻¹ (200 mg/dl) for 10–18 hours.
At the time when the determination was performed the average blood glucose was 16.2 mmol\textsuperscript{-1} (range 13.9–18.3 mmol \textsuperscript{-1}). Student's \( t \) test was employed in the statistical treatment using a 5\% limit of significance.

**Results**

**Blood Glucose, Body Weight, and Tibial Length**

At the time of the experiments blood glucose ranged between 16.9 and 28.0 mmol \textsuperscript{-1} (mean, 23.0 mmol \textsuperscript{-1}) in untreated diabetic rats. In the groups of rats with four weeks duration of diabetes the weight loss ranged between 50 and 130 g, the mean value being 91 g. After one, two, and five weeks duration of diabetes the mean weight loss was 47, 66, and 104 g, respectively.

Insulin-treated rats all survived and all gained weight. The weight increase after four weeks of treatment was 17 g (range, 5–35 g). In the group of two weeks of insulin treatment the mean weight increase was 12 g (range, 0–25 g).

Blood glucose values measured 24 hours after the daily insulin injection were 3.3, 4.9, and 7.9 mmol \textsuperscript{-1} when given by the 20th, 50th, and 80th percentiles. In other words blood glucose ranged between 3.3 and 7.9 mmol \textsuperscript{-1} 24 hours after the daily insulin injection in three of five days, the median being 4.9 mmol \textsuperscript{-1}. The corresponding percentile values of the two weeks treated group used in the neurophysiological experiments were 3.8, 5.4 and 8.2 mmol \textsuperscript{-1}.

The blood glucose variation during the 24 hour period in insulin treated rats with initial values ranging between 3.3 and 6.7 mmol \textsuperscript{-1} is illustrated in Fig. 1. The mean value of all measurements was 5.7 mmol \textsuperscript{-1} and the mean values of the various time intervals ranged between 3.4 and 8.8 mmol \textsuperscript{-1}.

The individual values of the measurements of tibial length in control and age-matched diabetic animals are shown in Fig. 2. Nearly identical values were obtained, 42.7 mm±0.7 (SD) and 42.5 mm±1.0, respectively. In the 48 weeks old control rats the length was 42.9 mm±0.8. This latter value is not significantly different from that of the 24 weeks old control rats.

**Light Microscopy**

It appears from Table 1 that the cross-sectional fascicular area and the endoneurial area still increased after the age of 24 weeks. This also applies to the mean fibre size and to the mean axon and myelin size.

The morphometric data of peripheral nerve after four weeks of diabetes are shown in Table 2. The mean fibre area and the axon area of myelinated fibres were decreased by 8.4\% and 13.4\%, respectively, the 2P-value being 0.039 and 0.00031. The endoneurial area and the endoneurial fraction were increased in diabetic rats, the 2P-values being 0.0016 and 0.000025, respectively.

The results of the insulin experiment are also
shown in Table 2. In the insulin-treated group mean fibre and axon size as well as the endoneurial area and the endoneurial fraction all differed significantly from the values of the untreated diabetic rats. None of these parameters differ significantly between controls and insulin-treated rats.

**NERVE CONDUCTION VELOCITY**

The results of the neurophysiological experiment are shown in Table 3. A gradual decline of nerve conduction velocity was observed as the duration of diabetes increased, the reduction being demonstrable already after one day of disease. After two weeks of diabetes the slowing of the nerve impulse was 4.2% (2P, 0.0089) and after five weeks 9% (2P, 0.0000015). The difference between the conduction velocities after two and five weeks duration of diabetes was also statistically significant, the 2P value being 0.0058.

Insulin treatment was able to prevent the impairment of peripheral nerve function. The conduction velocity for the group of insulin-treated diabetic rats did not differ significantly from the value of the control rats and was significantly faster than for the group of untreated diabetic rats (2P, 0.0051).

**ELECTRON MICROSCOPY**

*Myelinated nerve fibres* Within the test areas an average of 228 myelinated fibres was counted in each rat. The number of myelinated fibres of the common peroneal nerve was 1896±238 in the control group and 1798±204 in the diabetic group. The difference was not statistically significant.

The morphometric data of the electron microscopic analyses are given in Tables 4 and 5. Mean fibre size was 46.42 μm²±3.24 in the control group compared to 41.45 μm²±3.45 in the diabetic group. The difference in fibre size which amounts to 10.7% is statistically significant (2P, 0.0040).

In Table 5, 10 percentiles of individual cumulative size frequency distributions for the control group and for the diabetic group are shown. When identical percentiles are compared it appears that nerve fibres from diabetic rats are smaller for all percentiles, but the differences reached statistical significance only for the 90th and 99th percentiles.
Table 4  Morphometric data of cross-sectioned myelinated fibres from electron microscopic photomontages of part of common peroneal nerve in 10 diabetic rats and 10 control rats

<table>
<thead>
<tr>
<th>Fibre size (μm²)</th>
<th>Axon size (μm²)</th>
<th>Myelin size (μm²)</th>
<th>Slope coefficient (μ)</th>
<th>Y-axis intercept (μ)</th>
<th>Correlation coefficient (r)</th>
<th>SER fraction (×10⁴)</th>
<th>Mito-chondrial fraction (×10⁴)</th>
<th>Number of filaments (per 0.1 μm²)</th>
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*Significantly different from the control group.

Table 5  Fibre size for 10 percentiles of individual cumulative size-frequency distributions of electron micrographs of myelinated fibres in common peroneal nerve. Results are mean values for the diabetic group (n=10) and for age-matched control rats (n=10)

<table>
<thead>
<tr>
<th>Percentiles</th>
<th>Control group (μm²±2SD)</th>
<th>Diabetic group (μm²±2SD)</th>
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<td>10th</td>
<td>7.97±2.34</td>
<td>6.97±1.41</td>
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<td>20th</td>
<td>14.33±3.51</td>
<td>12.65±1.63</td>
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<td>21.39±3.71</td>
<td>19.29±2.28</td>
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<td>50th</td>
<td>39.53±6.15</td>
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<tr>
<td>60th</td>
<td>48.89±7.47</td>
<td>44.60±5.73</td>
</tr>
<tr>
<td>70th</td>
<td>62.32±12.25</td>
<td>55.22±7.06</td>
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<td>80th</td>
<td>75.80±11.80</td>
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<td>90th</td>
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<tr>
<td>99th</td>
<td>114.81±17.20</td>
<td>100.51±7.09*</td>
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</table>

*Significantly different from the control group.

The mean axon size (Table 4) was 18.61 μm²±1.72 in control rats as against 15.99 μm²±1.62 in diabetics. Thus the mean axon size of the diabetic group was reduced by 14.1%, the 2P value being 0.0026. The mean myelin size was reduced by 8.5% the control value being 27.81 μm²±2.73 and the diabetic value 25.46 μm²±2.50. However, the difference in myelin size was not statistically significant (2P, 0.060).

To obtain information about the relationship of the cross-sectional axon area to the cross-sectional myelin area, correlation analyses were performed for each animal of paired values of single fibres. The relationship of cross-sectional axon area to myelin area is decreased in diabetic rats when compared with age-matched controls. This can be seen from Table 4. The slope coefficient of individual regression lines was 0.570±0.085 in the control group as compared to 0.489±0.050 in the diabetic group (2P, 0.019). The mean regression lines for the two groups are shown in Fig. 3.
As to the morphometric analyses of the axon organelles, the SER fraction, the mitochondrial fraction, and the number of neurofilaments are given in Table 4. None of the differences were statistically significant. Furthermore, the number of SER profiles and mitochondrial profiles were found to be very similar in control and diabetic rats, 10.38±3.03 SER profiles per 100 μm² axon in control rats as against 12.42±4.83 in diabetics and 25.65±3.12 mitochondrial profiles per 100 μm² axon in control as against 26.76±2.42 in diabetic rats.

Unmyelinated nerve fibres The number of unmyelinated fibres in the common peroneal nerve was 3160±734 in the control group and 2864±1139 in the diabetic group. The mean fibre size was 0.468 μm²±0.066 in control as compared to 0.452 μm²±0.071 in diabetic rats. Neither these differences nor those of the calibre spectra were statistically significant.

As to the organelles, however, a pronounced reduction of 40% of the cross-sectional area fraction of the SER was found in the diabetic group. The SER fractional values were 0.0568±0.017 in control rats and 0.0343±0.013 in diabetics (2P, 0.0072). The number of SER profiles per 10 μm² axon counted in the controls and in diabetics were 14.60±4.43 and 13.60±4.08, respectively. Nearly identical values in the two groups were obtained for the mitochondrial fraction as well as for the number of mitochondrial profiles. The mitochondrial fraction in control and diabetic rats were 0.0421±0.0104 and 0.0459±0.0079, respectively, and the number of mitochondrial profiles per 10 μm² axon were 8.51±2.18 in control and 9.57±2.27 in diabetic rats. Individual values of fibre size, SER fraction, and the number of SER profiles are illustrated in Fig. 4.

![Fig. 4 Mean fibre size, fraction, and number of smooth endoplasmic reticulum profiles in each of eight control (○) and 10 diabetic rats (●).](image)

**Discussion**

It has been suggested that the reduction in myelinated nerve fibre calibre in diabetic rats is the result of a failure of growth including retarded skeletal length (Thomas and Sharma, 1976; Sharma et al., 1977; Jefferys et al., 1978). In previous papers we have demonstrated findings hardly compatible with this suggestion. First it was shown that control rats fed a restricted diet leading to a weight loss comparable to that of the diabetic animals have nerve fibres of normal calibre (Jakobsen, 1976a). Secondly, diabetic nerve fibres with reduced calibre were found to have normal nerve internodal length (Jakobsen, 1976b). Finally, the results of the present study show that tibial length is unchanged in our diabetic animals. We therefore conclude that the difference in calibre represents a true neuropathic change rather than a change caused by an ordinary retardation of growth in length.

![Fig. 5 Mean fibre size and standard errors at the beginning and end of the experiment in two groups of control rats (○), and at the end of the experiment in a group of diabetic rats (●).](image)

The electron microscopic part of the present study has confirmed our previous light microscopic finding of reduced calibre in streptozotocin diabetic rats, and the correlation analyses of the cross-sectional axon to myelin area of single fibres have given results in accordance with the previous findings of a decreased axon-myelin ratio (Jakobsen and Lundbæk, 1976). Moreover, it has been confirmed that all sizes of fibres seem to be thinner in diabetic rats (Jakobsen, 1976a).
How does the difference in fibre calibre arise? Figure 5 illustrates what has happened to fibre size in 28 weeks old diabetic and control rats after the four week experimental periods. If only the fibre size measurements were available it would have been tempting to conclude simply that the diabetic state had caused a relative shrinkage of the fibres or, in other words, had inhibited the increase in fibre calibre normally occurring in rats between the 24th and 28th week. However, the data available here on axons and myelin sheaths must also be taken into consideration.

A decreased axon-myelin relationship can be obtained in at least three different ways. Firstly, it may be the result of axonal shrinkage. Secondly, it can be caused either by an increase in the number of myelin lamellae or by an increased distance between the lamellae. A third possibility is that normal nerves of rats increase the axon size disproportionately to the myelin size between the age of 24 and 28 weeks while this does not occur in diabetic rats.

Whatever constitutes the precise mechanism behind the observed difference in fibre calibre and axon-myelin ratio, there can be no doubt about the observation that a structural axon abnormality is present in peripheral nerve in early streptozotocin diabetic rats. It is not decisive for this conclusion whether axonal shrinkage is involved or whether only inhibition of the normal increase in calibre during the experimental period takes place. The change in axon calibre is a prominent feature and may well explain the minor changes of the myelin sheath.

In myelinated axons no change of organelles was demonstrated with the applied microscopic and morphometric techniques. We do not conclude, of course, that the ultrastructure of organelles of myelinated axons is normal.

In unmyelinated fibres a 40% decrease of the amount of SER was observed in the diabetic rats. Since it is suggested that the compartment for the fast axoplasmic flow is vesicular elements, it might be that the SER decrease is a structural counterpart of a changed fast axonal transport within unmyelinated diabetic nerve fibres (Droz et al., 1975). A decreased transport of transmitter enzymes of myelinated fibres in sciatic nerve of streptozotocin rats with a diabetes duration of four weeks has been reported by Schmidt et al. (1975). The accumulation of acetylcholinesterase and choline acetylase proximal to a nerve crush was found to be reduced by 20% and 40%, respectively. Observations to be published from our own laboratory or axonal transport after injection of tritiated leucine and 14C-glucosamine into the fifth lumbar dorsal root ganglion of streptozotocin diabetic rats have shown abnormalities of the transport of glycoproteins.

It appears from the insulin experiments that the morphological and neurophysiological alterations observed in these studies can be prevented by insulin treatment leading to normalisation or near normalisation of the blood glucose level. Furthermore, the preventative effect of insulin in streptozotocin diabetic rats implies that the peripheral nerve changes are the result of diabetes, not of a toxic effect of streptozotocin itself.

Nerve fibre calibre increases with age in control rats and so does nerve conduction velocity (Miyoshi and Goto, 1973). Therefore, the effect of duration of diabetes on conduction must be examined in rats of identical age as in the present study. Decreased conduction velocity has been found in several studies of rats with a diabetes duration of from one week up to one year (Eliasson, 1964; Sharma and Thomas, 1974; Greene et al., 1975). Slowing of nerve conduction of large fibres was also demonstrated in the present experiment and, furthermore, a relation between the degree of slowing and the duration of the diabetic state was established. The decrease in conduction velocity developed gradually (Table 3) as did the difference in fibre calibre (Fig. 5). It is likely that at least part of the decrease in conduction velocity is explained by the difference in fibre calibre (Jakobsen, 1976a). Slowing of conduction by 3–4% could be demonstrated as early as 24–36 hours after the application of streptozotocin, at a time when blood glucose had been above 11.1 mmol 1−1 (200 mg/dl) for only 10–18 hours. This initial impairment of impulse propagation may well be explained by some other factors than difference in fibre calibre—for example, alterations of the axonal membrane at the node of Ranvier.

Finally, the relevance of the findings of the present study to human insulin deficiency diabetes requires discussion.

Segmental demyelination is observed regularly in histological studies of peripheral nerve from patients with several years of diabetes (Thomas and Lascelles, 1966; Chopra et al., 1969). Axonal loss is also present even though it has been claimed to be present particularly in the severe chronic cases (Thomas and Lascelles, 1966) or only in patients who have had diabetic neuropathy of longer duration and of insidious onset (Chopra et al., 1969). However, no conclusions about the interdependence of axon and myelin sheath changes can be drawn from the histological studies available on peripheral nerve changes in long-term diabetes.
As mentioned above, there is very little information about early structural changes in peripheral nerve of young patients with juvenile diabetes—that is, in a situation similar to that studied in our animal model. Bischoff (1973) mentions changes of axon organelles without the occurrence of segmental demyelination in a qualitative study of sural nerve biopsies from patients of various ages and with various types of diabetes. Very recently, he has reported results of another qualitative electron microscopic study of children and young patients who had had diabetes for less than one year (Bischoff, 1978). Severe axonal changes, again without the presence of segmental demyelination, were shown in this report. The results of quantitative analyses of these preparations will be of considerable interest.

At present it seems reasonable to assume that structural axonal changes occur very early in human beings with juvenile diabetes as in experimental insulin deficiency diabetes, that the axon abnormality is caused by the metabolic derangement characteristic of the diabetic state, and that segmental demyelination, absent in early juvenile diabetes and also in experimental diabetes (Sharma and Thomas, 1974; Jakobsen, 1976b) is a late secondary phenomenon.

References


Early and preventable changes of peripheral nerve structure and function in insulin-deficient diabetic rats.

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