



Myosin Va but not nNOS α is significantly reduced in jejunal musculomotor nerve terminals in diabetes mellitus

Arun Chaudhury^{1*}, Marcilio Hubner De Miranda-Neto², Renata Virginia Fernandes Pereira² and Jacqueline Nelisis Zanoni^{2*}

¹ Department of Surgery, Brigham and Women's Hospital, Harvard Medical School and VA Boston HealthCare System, West Roxbury, MA, USA

² Department of Morphological Sciences, Universidade Estadual de Maringá, Maringá, Brazil

Edited by:

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*Correspondence:

Arun Chaudhury, Department of
Surgery, Brigham and Women's
Hospital, Research 151, Room 2B103,
Building 3, 1400 VFW Parkway, West
Roxbury, MA 02132, USA

e-mail: arun_chaudhury@hms.
harvard.edu;

arunchaudhury.boston@gmail.com;

Jacqueline Nelisis Zanoni,

Department of Morphological
Sciences, Universidade Estadual de
Maringá, Maringá, Parana CEP
87020-900, Brazil

e-mail: jnzanoni@uem.br;

zanonijn@gmail.com

Nitric oxide (NO) mediated slow inhibitory junction potential and mechanical relaxation after electrical field stimulation (EFS) is impaired in diabetes mellitus. Externally added NO donor restore nitrergic function, indicating that this reduction result from diminution of NO synthesis within the pre-junctional nerve terminals. The present study aimed to investigate two specific aims that may potentially provide pathophysiological insights into diabetic nitrergic neuropathy. Specifically, alteration in nNOS α contents within jejunal nerve terminals and a local subcortical transporter myosin Va was tested 16 weeks after induction of diabetes by low dose streptozotocin (STZ) in male Wistar rats. The results show that diabetic rats, in contrast to vehicle treated animals, have: (a) nearly absent myosin Va expression in nerve terminals of axons innervating smooth muscles and (b) significant decrease of myosin Va in neuronal soma of myenteric plexus. In contrast, nNOS α staining in diabetic jejunum neuromuscular strips showed near intact expression in neuronal cell bodies. The space occupancy of nitrergic nerve fibers was comparable between groups. Normal concentration of nNOS α was visualized within a majority of nitrergic terminals in diabetes, suggesting intact axonal transport of nNOS α to distant nerve terminals. These results reveal the dissociation between presences of nNOS α in the nerve terminals but deficiency of its transporter myosin Va in the jejunum of diabetic rats. This significant observation of reduced motor protein myosin Va within jejunal nerve terminals may potentially explain impairment of pre-junctional NO synthesis during EFS of diabetic gut neuromuscular strips despite presence of the nitrergic synthetic enzyme nNOS α .

Keywords: axonal transport, myosin Va, nNOS, diabetes, neuropathy

INTRODUCTION

Slow inhibitory junction potentials (sIJP) are recorded using impaled electrodes within gastrointestinal smooth muscles in response to nitric oxide (NO) released from pre-junctional nerve terminals after electrical field stimulation (EFS). Independent studies have confirmed impairment of sIJP in muscle samples obtained from all portions of the gastrointestinal tract of animals with pharmacologically induced or genetically acquired diabetes mellitus (1–3). Mechanical studies have confirmed diminution of EFS-mediated nitrergic relaxations in diabetic enteric tissues like ileum and mid-colon (3–7). Some of these studies have also demonstrated restoration of nitrergic mediated responses in diabetic tissues with externally added NO donor to the organ bath (4–6), indicating that the diminution of nitrergic function in diabetic gut tissues resulted primarily from reduction or inhibition of NO synthesis within the pre-junctional nerve terminals.

Controversy exists in the reports regarding the content of neuronal nitric oxide synthase (nNOS) within the enteric nerve terminals in diabetic gut tissues. Limited data showing only a single low power microscopic field have been used to report reduced number of nitrergic axons traversing the neuromuscular wall in streptozotocin (STZ) induced diabetes (8). Other studies have reported

reduced dimer/monomer ratio of nNOS (9) or increase in total nNOS levels in diabetes (9, 10). However, the nNOS blots were run with extracts obtained from whole gut tissues, thus precluding specific information about nNOS contents within the nerve terminals *per se*. Studies have also reported normal nNOS enzymatic activity of diabetic whole gut extracts assayed *in vitro* (11). However, none of these studies provide unequivocal evidence about nNOS contents within the nerve terminals, the site of inhibitory enteric neuromuscular nitrergic neurotransmission.

Recently, evidence has been provided that mere presence of nNOS within nerve terminals is not adequate for pre-junctional NO synthesis (12–14). The regulation of nNOS within the nerve varicosities require multiple allosteric interactions, most notably, its positioning at PDZ-rich active zones that allow interfacing of water soluble nNOS α with membrane-bound palmitoyl-PSD95 (14, 15). Intriguingly, this binding of nNOS α is not stochastic and dependent on a Brownian kind of diffusion but rather relies on specific molecular interactions involving motor proteins like myosin Va that have the ability to deliver nNOS α to membrane-binding sites (12). Using a mouse model of myosin Va mutation, the dilute DBA/2J mice, it was shown that *in vitro* NO synthesis of enteric synaptosomes and NO-mediated sIJP and L-NAME

sensitive mechanical relaxations were impaired in gastric tissues of dilute mice (12, 14).

The purpose of the present study was to investigate two specific aims that may potentially provide pathophysiological insights into diabetic enteric nitrergic neuropathy: (a) is there any alteration in nNOS contents within enteric nerve terminals in diabetes (b) is there any alteration in myosin Va contents within enteric nerve terminals in diabetes. The first question was pursued to obtain unambiguous evidence about the state of nitrergic nerve terminals in diabetes at a fixed time point after diabetes induction. Based on preliminary evidence that local intravaricosity transport of nNOS α by myosin Va motor protein is important for efficient NO synthesis during neurotransmission (12, 13), we hypothesized that deficiency of myosin Va may contribute to impaired nitrergic neurotransmission in diabetes.

MATERIALS AND METHODS

INDUCTION OF DIABETES MELLITUS

All experimental procedures were conducted with approval from IACUC Committee of VA Boston HealthCare System (VABHS) and Committee of Ethics in Animal Experimentation from the Universidade Estadual de Maringá. Male Wistar rats (*Rattus norvegicus*) from the Central Biotery of Universidade Estadual de Maringá were utilized for the present investigation. Ninety-day-old rats were maintained in separate cages for 4 months prior to experimentation. After a 14-h fast, STZ (35 mg/kg body weight; Sigma, St. Louis, MO, USA) dissolved in freshly prepared 10 mM citrate buffer (pH 4.5) was intravenously injected. Animals with persistent plasma glucose >250 mg% were used for the study as a model of diabetes mellitus. Normoglycemic vehicle treated animals were used as controls. Jejunum were obtained after 4 months of diabetes induction. Details of tissue procurement and euthanasia procedures have been described earlier (16).

STAINING OF JEJUNAL WHOLE MOUNTS FOR MYOSIN Va AND nNOS α

Details of tissue fixation and staining were performed as per protocols described earlier (16). Jejunum was used as representative of gut neuromuscular strip. A polyclonal antibody raised against chicken brain myosin Va (17) and extensively characterized previously was used for myosin Va immunostaining (1 in 750 primary, prepared at University of São Paulo, Ribeirão Preto-campus). Tissue reaction was performed by incubating with hydrogen peroxide and diaminobenzidine (DAB). Omission of the primary antibody did not show any reaction product after labeling. nNOS α immunostaining was performed with a commercial antibody developed against epitope N₂₋₃₀₀ (H-299, 1 in 500, Santa Cruz Biotechnology) with appropriate negative control. Avidin-biotin histochemistry and Alexa Fluor fluorescent images were separately obtained to visualize nNOS α immunostaining. Uniform staining protocols were used.

LIGHT MICROSCOPIC IMAGING OF WHOLE MOUNTS AND ACQUISITION OF IMAGES

Digitized images were obtained with Zeiss Axioskop Plus light microscope at 200-fold magnification and stored for offline analyses. Same scaled images were used for comparative analyses, though digitally zoomed images may have been shown as representative figures for better depiction of the projected features.

PSEUDOCOLORING OF IMAGES FOR CREATING CONTRAST AND BETTER VISUALIZATION OF NEUROMUSCULAR STRUCTURES

Some image profiles were pseudocolored to create better contrast for visualization of neuromuscular apparatus. Images were opened in ImageJ (NIH freeware)¹ and converted to 16- or 32-bits prior to choosing a pseudocolor (sepia, magenta hot, blue orange icb, and glasbey) from the lookup table (LUT) for creating efficient contrast.

QUANTITATION OF SPACE OCCUPANCY OF NITRERGIC VARICOSITIES

Space occupancy of nitrergic varicosities was obtained using a simple derivative of graph theory. Graph theory relies on connectivity patterns based on number of vertices and edges and may be expressed as: $G = (V, E)$ (18, 19). The nerve varicosities may be inferred as the vertices (V) and the axonal processes (primary, as well as branches) to be edges (E). Based on the argument that cable length and sites of neurotransmission (junctions or varicosities) are critical for determining efficiency of neurotransmission (20), obtaining quantitative information on spatial densities will be a more reliable marker of neurites' arborization rather than random visualization of a snapshot of a whole mount microscopic field. Quantizing the pixel intensities of varicosities and the processes provide a semiquantitative approach toward estimating whether any changes occurred in the spatial density of nerve terminals after diabetes induction. In order to obtain pixel intensities of axonal processes, repeated thresholding was performed in ImageJ to remove background intensity. The auto-threshold function (Huang, intermodes, Li, maximum entropy, and isodata) in ImageJ was used for these analyses. RGB images were converted to 8 or 16-bit prior to thresholding.

3D SURFACE MODELING TO VISUALIZE MYOSIN Va MUSCULOMOTOR TERMINALS

The plugin 3D interactive surface plot in ImageJ was used to visualize the stained varicosities and myenteric plexus. Unaltered images were used for these quantitative analyses. The spike height and color hue are representative of immunoexpression intensity. Same parameters of grid size, smoothing, perspective, and lighting were used for all whole mounts analyzed.

ESTIMATION OF nNOS α CONTENTS WITHIN NITRERGIC VARICOSITIES

Varicosities were carefully outlined with freehand selection tool in ImageJ and area and integrated densities estimated for nNOS α concentrations within varicosities and intensities normalized to area. Axonal processes were carefully traced from the cell body toward the ramification on the muscular wall and followed till the process ended or faded in a different plane. This information was also used to map a static kymograph that provided insights into nNOS α axonal transport in vehicle treated and STZ-induced diabetes. Linear distances between varicosities were measured. Raw images were used for intensity analyses.

NEURONAL PROCESS TRACING

Free hand tools were used to trace nitrergic neurons on planar unscaled whole mounts in MS Paint. The trace was independently

¹<http://imagej.nih.gov/ij>

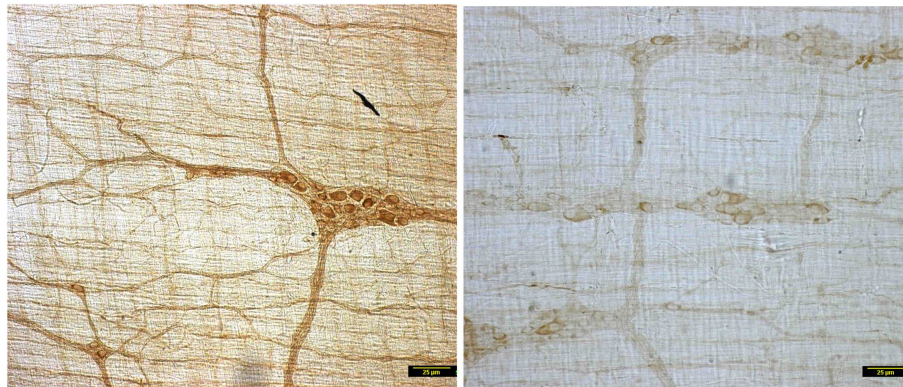


FIGURE 1 | Low power micrographs from jejunal whole mounts of vehicle treated (left) and streptozotocin induced diabetic rats (right). Note the obvious differences in DAB staining intensity of myosin Va between the two panels. Scale bar, 25 μ m.

visually verified in Neuron² (data not shown) (21). Random selections of neurons were made from independent whole mounts obtained from different rats.

STATISTICAL ANALYSES

Quantitative data was expressed as mean \pm SEM. Two-tailed *t* test was used to compare difference between means of the parametric datasets.

RESULTS

MYOSIN Va IMMUNOREACTIVITY IS SCANT OR NEARLY ABSENT IN THE NEURO-SMOOTH MUSCLE NERVE TERMINALS OF JEJUNUM

In contrast to the vehicle treated rats, induction of diabetes by STZ resulted in nearly complete loss of myosin Va within the nerve terminals of the neuronal processes ramifying on the muscularis externa (Figure 1). Whole mounts of jejunum stained with myosin Va specific antibody failed to show the brown reaction product of DAB staining that was visualized in the enteric tissues of normal rats (Figure 1). The absence or paucity of myosin Va staining overlying the muscularis externa was apparent during light microscopic imaging.

MYOSIN Va IMMUNOREACTIVITY IS VARIABLY DECREASED OR NEARLY ABSENT IN MANY NEURONAL CELL BODIES OF MYENTERIC PLEXUS OF JEJUNUM

In wild type condition, myosin Va immunoreactivity was strongly present in the cell bodies of all neurons present within the myenteric plexus. The brown reaction product was well visualized and uniformly distributed across the whole cytosol of the neuronal soma. Myosin Va immunoreactivity was also well visualized in the axon initial segment (AIS) of the ganglionic neurons and extended into the processes that link different cells within the ganglion, as well as axonal extensions into the muscularis (Figure 2). There was diffuse staining for some length of the initial parts of the axonal extensions, but thereafter, myosin Va immunoreactivity was seen as regular punctate appearance, indicating probable concentration

of myosin Va in the nerve varicosities of the neuro-smooth muscle neuromuscular junctions.

In contrast, myosin Va immunoreactivity was severely diminished in most neuronal cell bodies in diabetic jejunal whole mounts (Figure 2). In the neuronal cell bodies that demonstrated reduced staining, myosin Va immunoreactivity did not extend even into the AIS (Figure 2). Significantly, no myosin Va immunoreactivity was seen even in the initial portions of the axonal extensions in diabetic jejunum. Myosin Va immunoreactivity was only scantily and rarely present in the nerve varicosities overlying the muscularis in diabetic tissues (Figure 3). Light microscopic imaging reveals the variance of the myosin Va staining and distribution within the ganglion neurons and its arborizations in diabetic tissues (Figure 4). Molecular techniques like western blotting of whole tissue protein extracts would fall short in identifying this significant observation. Because of only sporadic presence of spotty myosin Va-IR within nerve varicosities of diabetic tissues (Figure 3), quantitative myosin Va concentrations within nerve terminals of the neuromuscular junctions were not obtained.

UNLIKE MYOSIN Va, nNOS α IMMUNOREACTIVITY IS NORMAL IN THE NEURO-SMOOTH MUSCLE NERVE TERMINALS OF JEJUNUM

nNOS α immunoreactivity within the nerve terminals in vehicle treated and STZ-induced diabetic jejunal tissues was quantitatively examined. nNOS α was present within the nerve terminals of both vehicle treated and STZ-treated rats (left panels of Figures 5 and 6). Also note that the nerve terminals and nitrergic axons form loop like structures, reminiscent of Kirchoff-type circuits. This distribution may serve to propagate inhibitory nerve signals simultaneously over a segment of muscle, though no electrophysiological evidence or imaging evidence of simultaneous NO production over the loop with a single electrical stimulus has ever been obtained.

nNOS α IMMUNOREACTIVITY IS NORMAL OR VARIABLY DECREASED IN THE NEURONAL CELL BODIES OF MYENTERIC PLEXUS OF JEJUNUM

The variance of nNOS α expression is demonstrated in the right panels of Figures 5 and 6 (vehicle and STZ treatment, respectively). The cellular expression of nNOS α is comparable between vehicle

²<http://www.imagescience.org/meijering/software/neuronj/>



FIGURE 2 | Myosin Va immunoreactivity extending from the cell body to the processes of Dogiel type 2 neurons emanating from myenteric plexus. In vehicle treated group (left panel), myosin Va is present in the initial segment, as well as axonal processes extending to the muscle layers. Myosin Va is also present in the interneuronal processes within the

ganglion. In contrast, note that in the diabetic rats, despite the presence of secondary plexus linking the ganglia and tertiary extensions of nerve processes into the muscle layers, the brown DAB reaction product representative of myosin Va presences is scant or nearly absent in these processes. Scale bar, 25 μm .

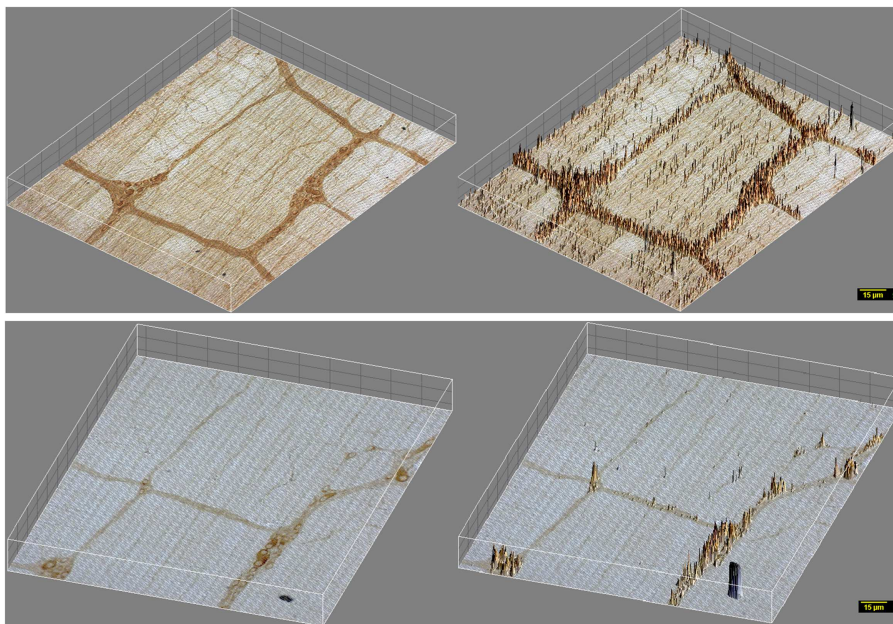


FIGURE 3 | 3D surface plot of DAB staining intensity of myosin Va in WT and diabetic jejunum. Note that in vehicle treated rats (upper panels), staining spikes present in the muscle segments interposed between the ganglia and plexuses have dense distribution of myosin Va spikes in the nerve terminals, while these spikes within the muscularis are almost absent in

diabetic tissues (lower panels). These nerve terminals are the sites of neuro-smooth muscle neurotransmission. Note the lighter hue of the staining spikes within the neuronal cell bodies, representative of reduction in myosin Va genomic expression within the neuronal soma in diabetic tissues in comparison to vehicle treated rats. Scale bar, 15 μm .

treated normal (Figure 5) and STZ-treated diabetic jejunum (Figure 6). While most of the cells are uniformly stained in the wild type tissues with comparable pixels of expression, the fourth segment in the right panel (Figure 5) shows cell bodies with

comparatively low expression of staining. The reason to demonstrate this variance is to stress that mere comparative expression from whole tissue extracts by protein blotting techniques is not optimal to obtain evidence regarding the neuropathology

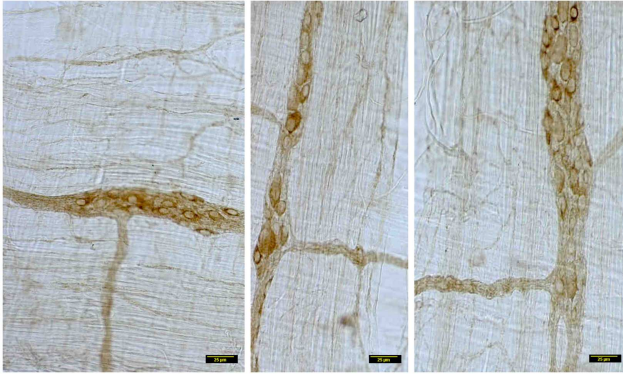


FIGURE 4 | Details of myosin Va staining of ganglionic neurons and their processes in diabetic jejunum. Note the heterogeneity of appearance of myosin Va in the cell bodies of the neurons in the myenteric plexus. In the left panel, note the moderate staining in the cell bodies of many neurons. Also note that the secondary plexus running to the left of the image has some brown DAB staining. Some of the neuromuscular varicosities have faint brown staining. In the middle panel, note the enlarged neuronal soma with unhealthy appearing nucleus, possibly indicating a degenerating neuronal cell body. The increase in size may be related to the cellular accumulation of myosin Va, possibly resulting from stasis of myosin Va axonal transport. Also note the absence of brown staining in all of the neuronal processes ramifying through the muscular wall. The right panel recapitulates information represented in the middle panel and in addition, shows a full range of neuronal cytopathology. Note that in the upper pole, two neurons do not have any cytoplasmic staining and the lower neuron is possible degenerative, as evidenced by unusual nuclear appearance. Note very scant or nearly absent staining in the interneuronal processes, secondary plexi as well as tertiary neuromuscular nerve varicosities. Also note that some cells (top right, also one cell body in the right on left panel) show intense dark brown staining. This heterogeneous appearance may result from enhanced myosin Va transcription but may have also resulted from myosin Va aggregates formation. Scale bar, 25 μ m.

of nNOS α expression. Some of the neuronal cell bodies demonstrate lower levels of expression of nNOS α (Figure 6). However, nNOS α was present in the neuronal processes arising from these cell bodies. Most whole mounts actually demonstrated normal levels of expression of nNOS α in the diabetic tissues. Occasionally, apoptotic neuronal cells with high nucleocytoplasmic ratios and skeletonized outlines were observed (Figure 6, lower segment of the right panel).

NITRERGIC NEURONAL PROCESSES ON MUSCULARIS EXTERNA ARBORIZE NORMALLY AND SHOW NO RETRACTION IN DIABETIC JEJUNUM

Observations of the whole mounts show wide variance in the distribution patterns of the processes emanating from the ganglionic neuronal cell bodies and extending to the muscularis externa. Representative tracings indicating this variance is demonstrated in both vehicle treated and diabetic tissues in Figure 7. Several nitrergic neuronal processes were traced and compared between vehicle treated and diabetic tissues. The neuronal arborizations were extensive and cable length comparable between sham treated and diabetic tissues, indicating no retraction of neuronal processes of nitrergic neurons that ramify on the muscularis in diabetes (Figure 7, lower panel). Semiquantitative parameters like the

distance of the farthest point on the axonal process from the centroid of the neuronal cell body were used to compare neural arborization. These data were comparable between vehicle and STZ treatment (data not shown).

nNOS α IS NORMALLY DISTRIBUTED IN THE CELL BODIES OF MYENTERIC PLEXUS AND EXTENDS INTO THE DENDRITES OF DOGIEL TYPE 1 NEURONS AND INTERNEURONAL PROCESSES WITHIN GANGLIA IN DIABETES

nNOS α is present not only in the nerve terminals in diabetes, but also normally distributed within the cell bodies as well as the dendrites of Dogiel type 1 neurons. Images were converted to 16-bits and pseudocolor coded prior to thresholding for optimal visualization of the neuronal cell body and processes. The distribution of nNOS α in wild type and diabetes in the proximal processes were comparable (Figure 8). Interneuronal processes between smooth Dogiel type 2 neurons and axonal extensions into the muscle contained comparable levels of nNOS α in vehicle treated and diabetic tissues. These qualitative observations indicate that at the time period examined, nNOS α axonal and dendritic transport is not impeded despite diabetic condition of the rats. Again, it may be appreciated that the genomic expression of nNOS α is variable and may be slightly diminished in some ganglionic neurons. The intact axonal transport of nNOS α in diabetes is sharply contrasted against the nearly absent axonal transport of myosin Va in diabetes at the specific time point examined.

SPACE OCCUPANCY OF NITRERGIC VARICOSITIES ARE COMPARABLE BETWEEN VEHICLE TREATED AND STZ-INDUCED DIABETIC JEJUNUM

Because of the variance in the distribution patterns of nitrergic nerve processes within the muscular wall, examination of a single field of whole mount may not be accurately reflective of axonal arborization. In order to obtain quantitative evidence of intactness of arborization pattern of nitrergic axons and varicosities, we incorporated a simple application of graph theory. Essentially, we argued that a unit of muscle segment delimited on the sides by ganglion and the primary and secondary plexus constitutes a unit of muscle bed that is supplied by axonal processes extending into this region from the adjacent plexi and cell bodies. One short coming of this approach is that only a planar layer was examined, while it is possible that the ramification of the nerve fibers may be deep to the visualized tissue plane. Considering obtaining a semiquantitative estimate, only planar images were considered, ignoring the stereological aspect. In comparison to previous reports, which have shown a single section to comment on nitrergic nerve process distribution (8), we estimated the total intensity of the axonal extensions and normalized to the muscular area of distribution. The total intensity was obtained by numerous iterations of thresholding (Figure 9B), the averaging technique contributing to closely accurate estimates of pixel integrated densities. In this way, we computed the space occupancy of nitrergic processes. Because the intervaricose segments are much less stained than the varicosity regions, so it is logical to extrapolate this calculation to be an estimate of space occupancy of nitrergic varicosities. Though some diabetic sections showed a diminution in extension of nitrergic processes across muscle bed (lowest

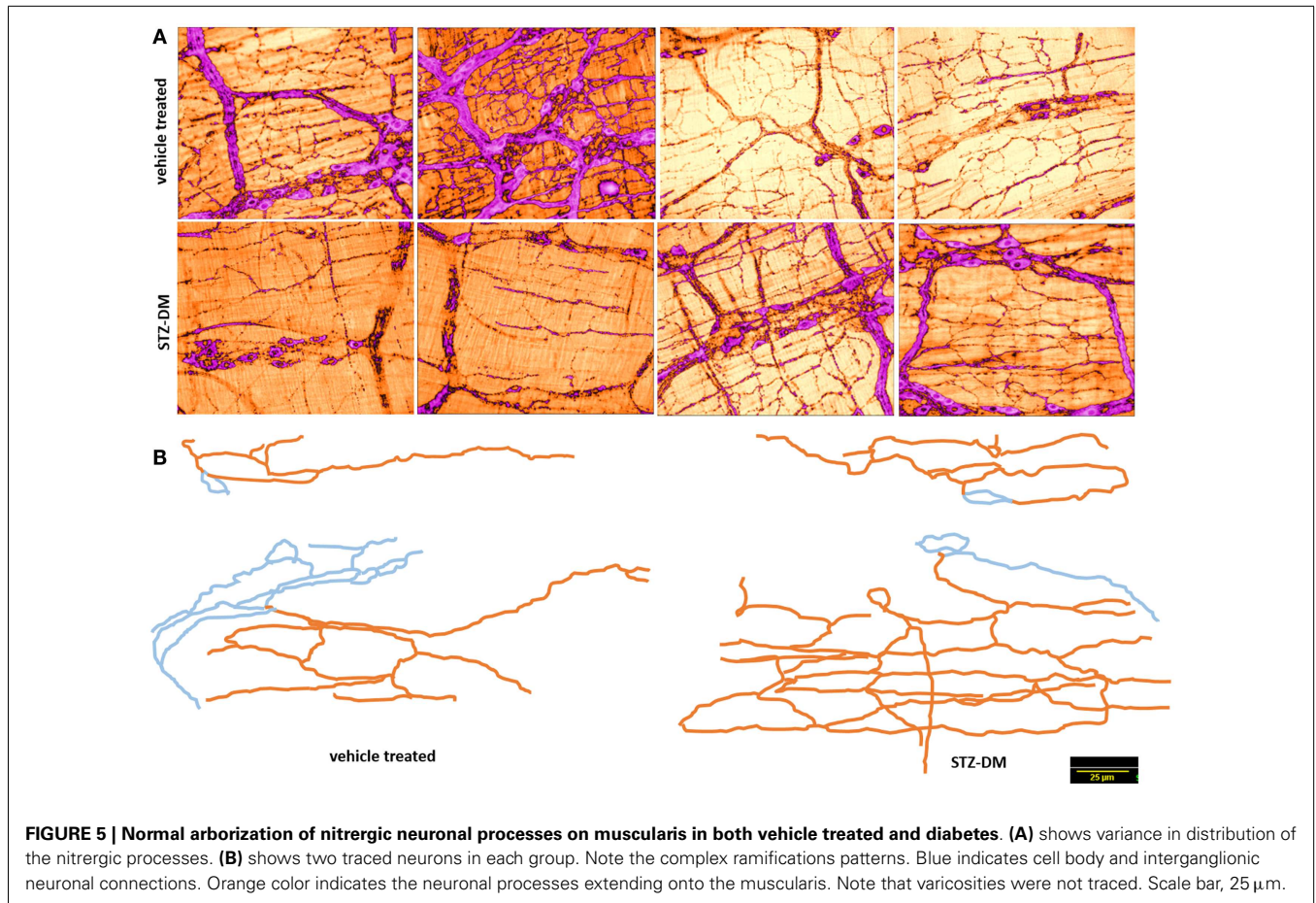


FIGURE 5 | Normal arborization of nitrenergic neuronal processes on muscularis in both vehicle treated and diabetes. (A) shows variance in distribution of the nitrenergic processes. **(B)** shows two traced neurons in each group. Note the complex ramifications patterns. Blue indicates cell body and interganglionic neuronal connections. Orange color indicates the neuronal processes extending onto the muscularis. Note that varicosities were not traced. Scale bar, 25 μm .

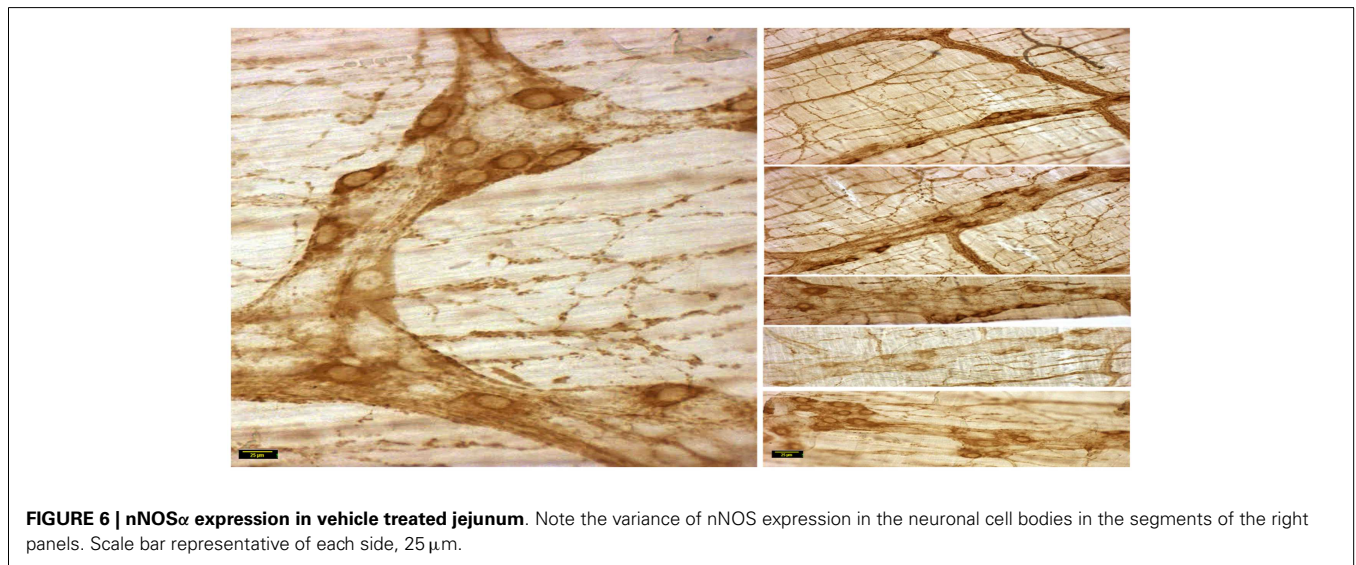


FIGURE 6 | nNOS α expression in vehicle treated jejunum. Note the variance of nNOS expression in the neuronal cell bodies in the segments of the right panels. Scale bar representative of each side, 25 μm .

count 153 $\text{AU}/\mu\text{m}^2$), cumulative analyses show comparable levels of the space occupancy of nitrenergic varicosities (215.25 ± 0.74 vs. $209.84 \pm 0.82 \text{ AU}/\mu\text{m}^2$, $p = 0.42$, two-tailed t test) (Figures 9A and 9C).

nNOS α CONCENTRATIONS ARE COMPARABLE BETWEEN VEHICLE TREATED AND DIABETIC MUSCULOMOTOR NERVE TERMINALS

The preceding analyses suggest that while myosin Va expression is nearly absent in the motor nerve terminals and significantly

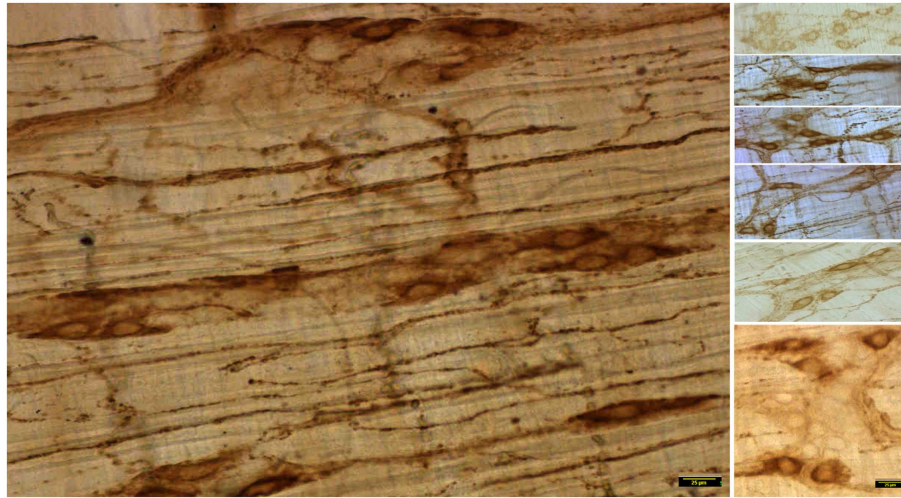


FIGURE 7 | nNOS α expression in STZ-treated diabetic jejunum. Note the variance of nNOS expression in the neuronal cell bodies in the segments of the right panels. Note reduction of nNOS α expression in some cell bodies.

However, the neuronal processes emanating from these cell bodies show nNOS α expression. Note that the lowest panel shows some degenerating neuronal cell bodies. Scale bar representative of each side, 25 μ m.

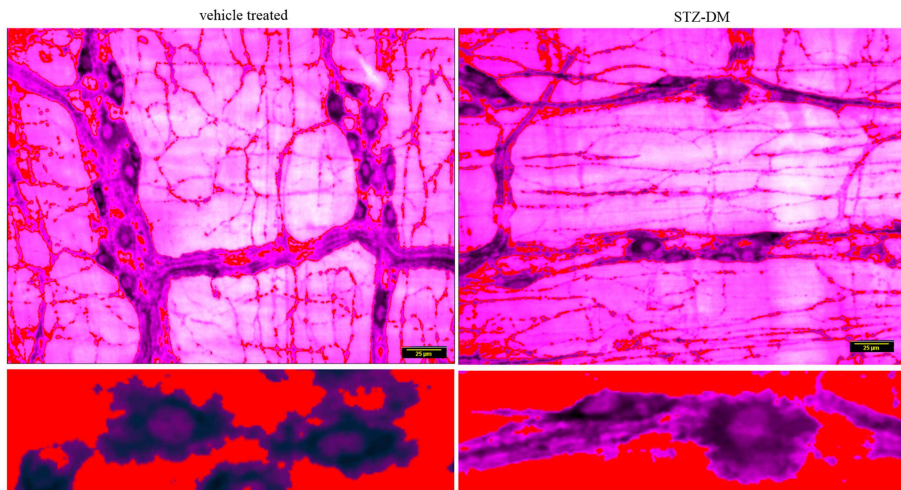


FIGURE 8 | Normal staining of nNOS within dendritic processes of Dogiel type 1 neurons and interganglionic neuronal processes in both vehicle treated and STZ-induced diabetic jejunum. The lower panels show a digitally zoomed image showing that nNOS extends into

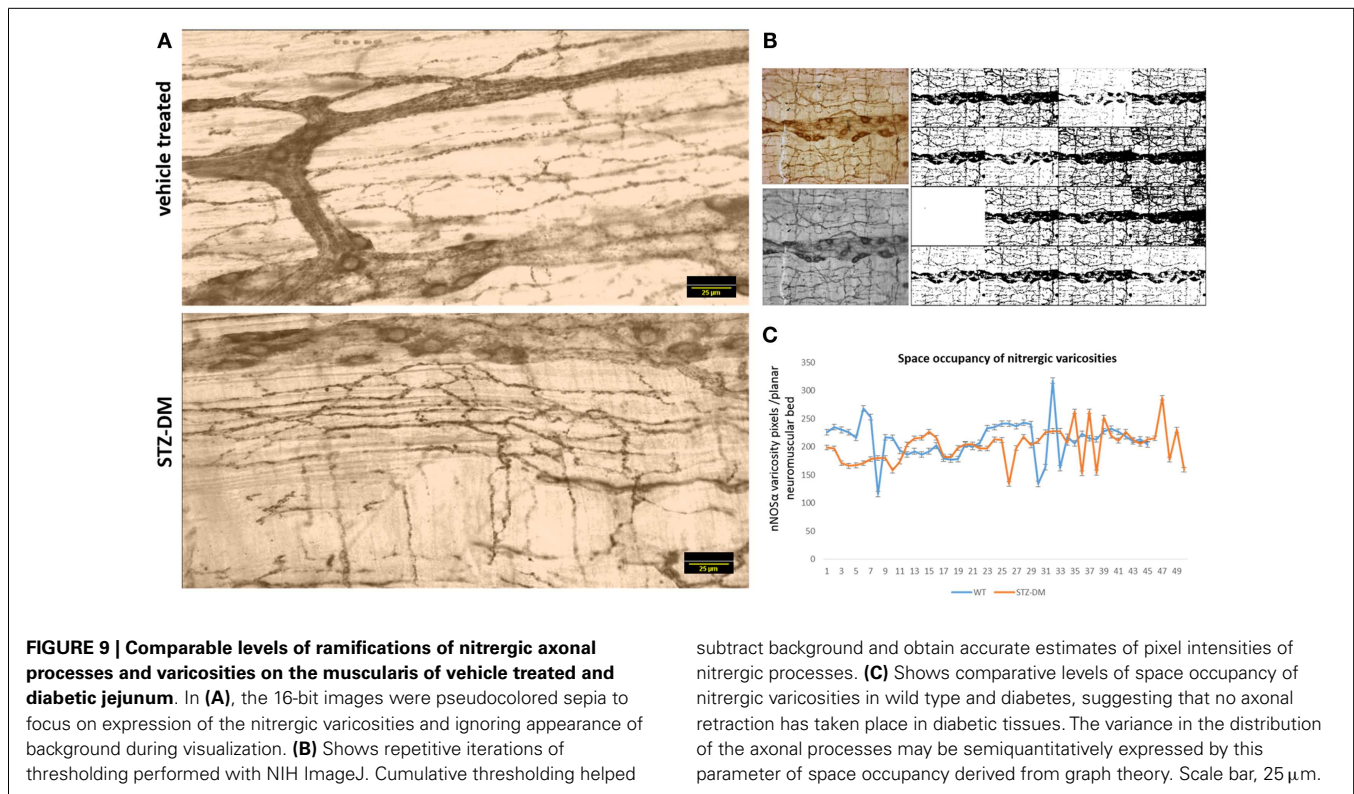
the proximal processes of the nitrergic neurons in both wild type and diabetic jejunal tissues. Renyi entropy type thresholding of pseudocolored images were used to observe the neuronal branch pattern morphology. Scale bar, 25 μ m.

reduced or absent in the pan-neuronal soma of the myenteric plexus from which the processes arise, nNOS α -positive nerve fibers appear normal in diabetic jejunum. Fluorescent images of nNOS α were used to examine specific nNOS α expression within the nerve terminals (Figure 10A). The images were thresholded to define the shape of individual varicosities and ease in estimation of the area of the varicosity (Figure 10B). Fluorescence intensities of nNOS expression were normalized to the areas and these values were comparable between vehicle treated and STZ-induced diabetic jejunum (18.9 ± 0.24 vs. 17.32 ± 0.15 AU/ μ m², $p = 0.8$, two-tailed t test). The intervaricose distances between nitrergic

varicosities were comparable between vehicle treated and STZ-induced diabetes (3.22 ± 0.27 vs. 3.08 ± 0.18 μ m) (Figure 10C).

STATIC KYMOGRAPH SHOWS INTACT nNOS α AXONAL TRANSPORT IN DIABETIC JEJUNUM

Static kymograph of nNOS α axonal transport was computed (Figure 10C). Nine randomly chosen nitrergic neurons from different rats with traceable and distinctly appearing varicosities in each group were used to obtain quantitative estimate of nNOS α axonal transport. The kymograph shows that despite some reduction of nNOS α expression in few of the nerve terminals, nNOS α



distribution is comparable between wild type and diabetic axonal processes, and that nNOS α reaches the farthest neuromuscular terminals in diabetic rats. This indicates that the axonal transport of nNOS α is likely not impeded in diabetes, unlike myosin Va axonal transport, which is severely impaired.

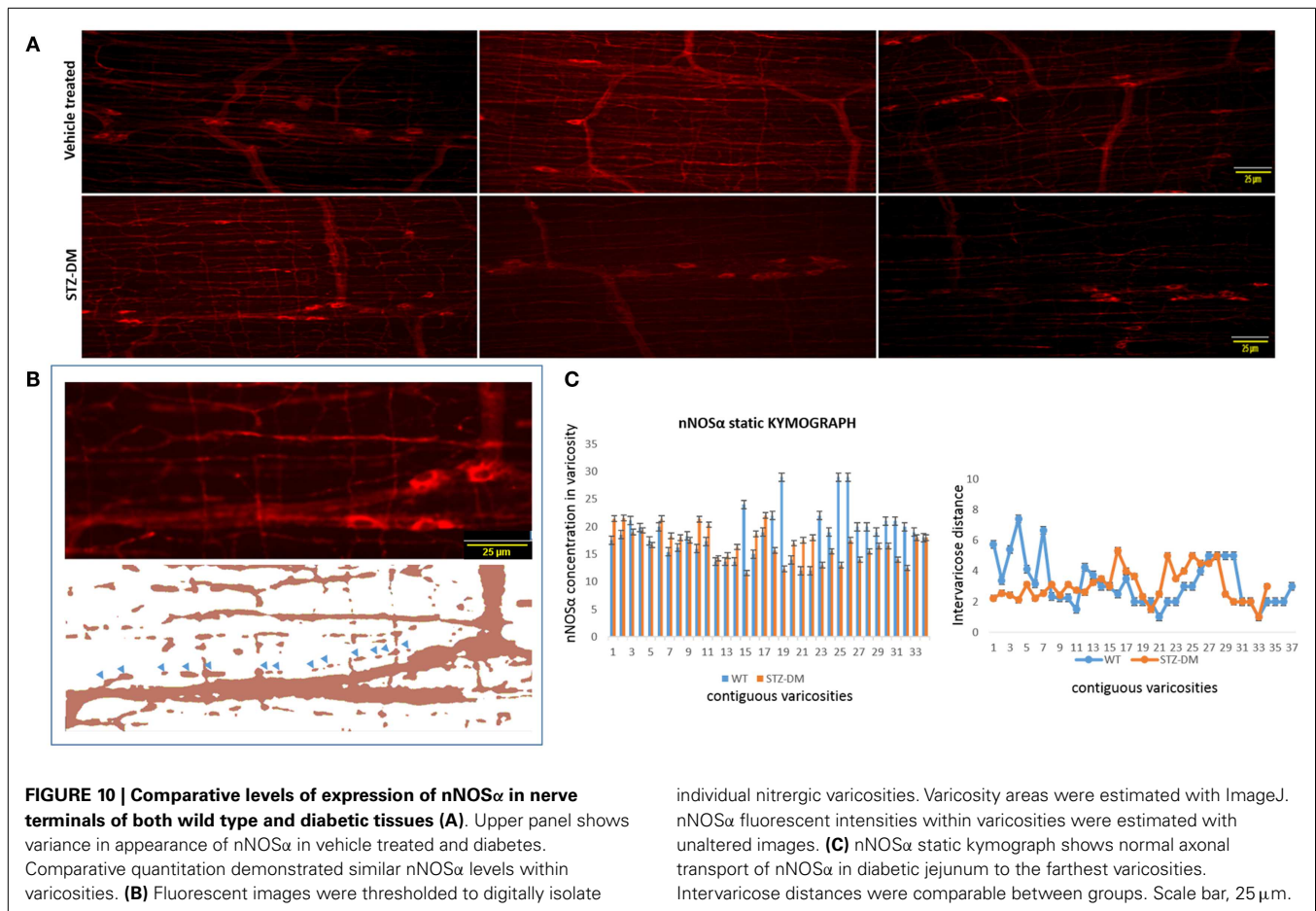
DISCUSSION

The results of the current study reveal that in contrast to vehicle treated rats, jejunum obtained 16 weeks after rats were injected with low dose STZ show: (a) nearly absent myosin Va expression in nerve terminals of fibers traversing the enteric neuromuscular bed, (b) variable decrease of myosin Va in cell bodies within the myenteric plexus, with near absence of myosin Va in several ganglionic neurons, and (c) some presence of myosin Va in the AIS of ganglionic cell bodies that stained for myosin Va, but near complete absence of myosin Va staining even in the initial segments of the primary plexus. In contrast, nNOS α staining in diabetic jejunum neuromuscular strips showed: (a) near intact expression of nNOS α in neuronal cell bodies, with reduction of expression in few cell bodies, (b) intact presence of nitrergic nerve fibers, with normal ramification and arborization patterns of nitrergic nerve fibers, normal density of nitrergic nerve fibers and terminals in comparison with untreated animals and normal concentration of nNOS α within a majority of nerve terminals and intact axonal transport of nNOS α to distant nerve terminals.

This report has examined the distribution of nNOS α quantitatively in diabetic enteric nerve terminals. While there was reduction of nNOS α expression in some cell bodies, as well as in some nerve terminals, the images, as well as the quantitative

analyses show that there is a wide variance of expression of nNOS α both in wild type tissues, as well as after STZ treatment. Thus, mere western blots of whole gut extracts (9, 10) may not be optimal approach for revealing the dynamic changes in nNOS α expression. Cumulative analyses, in fact, reveal that there is no statistically significant reduction of nNOS α expression within the nerve terminals in STZ-treated rat jejunum. Studies have demonstrated and discussed the progressive nature of diabetic autonomic neurodegenerative changes (3, 22). The results presented in this study needs to be cautiously interpreted in the background of the single time point that has been examined.

The findings of the unaltered expression of nNOS α within the nerve terminals are sharply contrasted with the near absence of myosin Va within the neuromuscular nerve varicosities. Quantitative density plots revealed only patchy and spotty appearance of myosin Va-positive nerve terminals after STZ treatment. It may be noted that there is a wide variance of myosin Va expression in the cell bodies comprising the ganglia in diabetes. A wide range of changes has occurred in myosin Va expression within the neuronal soma after diabetes induction. In some cell bodies, the myosin Va expression appears comparable to those of untreated animals. However, most neurons show moderate to severe reduction of expression of myosin Va. Interestingly, even when myosin Va expression is visualized within the neuronal soma, it is very difficult to observe any amount of expression of myosin Va within the processes emanating from the Dogiel type 1 and 2 neurons. One possibility may be that the primary processes may be in a different plane than the one visualized and imaged. However, close



visualization failed to show myosin Va in the axonal extensions. There may be two plausible reasons, which may have contributed to this state: (a) the low synthesis of myosin Va may have produced inadequate levels of the protein so that no substantial protein was available for transport to the axon, (b) this observation may also result from alterations in the protein quality, e.g., glycosylation of myosin Va may have transformed the protein to an effete nature, precluding its normal function and transit to the AIS. Though myosin Va may be normally partially glycosylated, other studies have shown that excessively glycosylated myosins have reduced ATPase activity and ability of force generation through actomyosin formation (23–25). Succinylated wheat germ agglutinin (sWGA) based assays have shown that nNOS is not glycosylated (26), thus creating a dilemma of why nNOS function is so severely affected by the hyperglycemia of diabetes. The results of the current study provide the first possible hints into the molecular pathophysiology of how nitrergic dysfunction may occur in diabetic nerve terminals due to deficits of molecular motors.

The near complete absence of myosin Va within the proximal portions of neuronal extensions may also have resulted from defects in spectrin, which may form an unusual barrier at the axon hillock and prevented protein migration (27). This possibility is unlikely, since the results suggest that transport of other soluble proteins like that of nNOS α has not been impeded. Yet another

possibility may be the result of defect of an allosteric partner for myosin Va that participates in axonal transport of myosin Va. Some preliminary evidence has been provided that neurofilament cooperatively participates in axonal transfer of myosin Va (28). It has been reported that neurofilament levels may be affected in pre-diabetic condition (29) and may decrease very early in diabetes (30). Decreases in neurofilaments have been reported in myenteric neuronal processes in high-fat diet induced diabetes (31). Earlier, it has also been demonstrated that despite the normal presence of neurofilaments in the soma and axons of myenteric neurons, a RT97 specific phosphorylated neurofilament epitope was deficient in diabetic rat ileum, and this was reversed when strips were incubated in insulin for 2 h (32). Alterations of neurofilaments may have potentially contributed to the total stasis of myosin Va within the nerve cell bodies seen in the current study. Electron micrographic examinations of neuronal processes within stomach tissues obtained from patients with diabetic gastroparesis have shown derangement of orientation of neurofilaments (33). Whether such spatial alterations of neurofilaments contribute to pathophysiology of loss of axonal transport of myosin Va in diabetic myenteric plexi remains to be examined.

The current investigation provides insights into axonal transport of nNOS α and some of the early changes seen in diabetic nerve

cell bodies and processes. Overall analyses show that nNOS α is present in the farthest varicosity away from the cell body, in comparable concentrations to those seen in untreated animals. The arborization patterns remain unaltered and the numbers of nerve terminals on each process are comparable between untreated and diabetic samples, indicating that there is no neurite retraction. Earlier studies have reported loss of nitrergic neuronal cell bodies even in the early phase of STZ-induced diabetes (16, 31, 34–36), though not all studies examining at similar time points (8–16 weeks) after STZ treatment found loss of myenteric neurons (8, 37). Despite probable neuronal loss, the current study demonstrated intactness of spatial arborization of nitrergic nerve terminals in the jejunum after 16 weeks of diabetes induction. Our results are in coherence with earlier observations of intactness of nNOS expression made in stomach biopsies obtained from patient samples of diabetic gastroparesis (38). Within the cell bodies, we have demonstrated comparable levels of expression of nNOS α between untreated and diabetes. The montage images of the cell bodies show the wide variability of expression, reiterating that mere protein estimates after blotting whole tissue extracts will provide inadequate insights into the state of expression of nNOS α in untreated and diabetic tissues. The results of the current study show that density of the enteric nerve terminals serving unit area of muscularis externa is comparable between diabetes and untreated, as also the intensity of nNOS α within individual varicosities. Some diabetic tissues have shown reduced expression of nNOS α both within the cell bodies and the varicosities along the axons extending from these cell bodies, but cumulatively, there were no significant alterations. An earlier case report demonstrated loss of nitrergic innervation in a human jejunum sample obtained from a patient with long standing diabetes (39). These divergences in findings may have primarily resulted from ongoing progressive pathophysiology of nitrergic neurotransmission system.

The dissociation between unaltered expression of nNOS α and grossly reduced expression of myosin Va within enteric neuromuscular nerve terminals is the most significant finding of this study. This finding is highly significant, and provides a possible basis of explanation why repeated observations have been made of impaired pre-junctional NO synthesis, despite presence of NO synthase within nerve terminals (1, 4, 10, 11). Because intravaricosity myosin Va deficiency has been demonstrated to be critical for NO synthesis during neurotransmission (12, 14), it may be reasonable to speculate that the findings of deficiency of myosin Va within diabetic enteric nerve terminals may explain the impaired pre-junctional NO synthesis seen in diabetes. It has been reported that altered nNOS dimer/monomer ratio may be the basis of impaired NO synthesis (9), and this may have resulted from deficiency of tetrahydrobiopterin (40). Detailed biochemical analyses have revealed that tetrahydrobiopterin may contribute to dimer stabilization rather than dimer formation (41). It is possible that myosin Va deficiency results in insufficient delivery of nNOS α to the varicosity membrane, the likely function of myosin Va within varicosities (12, 42). Future studies comparing membrane-bound nNOS α dimer in wild type and diabetic tissues may resolve these issues.

The present study probably is the first analyses examining axonal transport of nNOS α and its force-generating cytoskeletal

allosteric partner myosin Va and provides valuable insights into the mechanisms of axonal transport of soluble proteins. For myosin Va in the untreated group, myosin Va was visualized as discrete puncta along the nerve processes traversing the muscularis. A continuous haze of staining was only visualized in some initial segments within the primary plexus. Being a soluble protein, myosin Va likely undergoes slow transport, but may piggyback onto faster transport systems as it progresses into the axon. Importantly, there must be some mechanism for its capture at the axonal varicosities, and this possibly contributes to the punctate appearance farther along the axons. The intervaricose region here remains myosin Va-stain free, indicating that myosin Va is possibly transported as a macromolecular assembly along the axon. In diverse neuronal systems, actin and NF-L have been reported to transport myosin Va (28, 43). It remains to be examined, which allosteric protein partner(s) contribute to axonal transport of myosin Va within enteric neuronal processes. Interestingly, the axonal transport of nNOS α is not affected or only minimally affected at the 16-week time point examined in the present investigation. This may have resulted from intact transport mechanism(s) of nNOS α during the initial weeks of diabetes induction.

Most likely, nNOS-myosin Va do not form complex during transport; otherwise, nNOS α stasis would also have occurred in the neuronal cell bodies like that of myosin Va. This argument is brought in because myosin Va and nNOS may be linked by the common light chain LC8 (12, 44). However, it may be inferred from the current observations that myosin Va and nNOS are independently transported within the myenteric axons and tertiary processes. Recent evidence and debates are revealing that soluble proteins may not undergo the true form of slow axonal transport as originally reported for soluble proteins with *in vitro* experiments and radioactive pulse chase labeling (45–47). Such slow rates of about 1 mm a day are practically impossible to deliver proteins at the processes located several orders of distances away (48–51). Incipient but convincing evidence have demonstrated that soluble proteins tag on fast axonal transport systems and gets rapidly transported to distal nerve terminals (52). The major fast transport systems involve microtubules and kinesins (53). Specific nNOS–kinesin interactions have not been reported, but kinesin function deficit resulting from altered GDNF signaling in KIF26A knockout has been reported to cause Hirschsprung-like phenotype with impaired nitrergic function (54). This important issue merits future investigations. Though fast axonal transport is also affected in diabetes, this usually results from long duration of hyperglycemia. How nNOS α axonal transport is affected as a result of prolonged duration of diabetes is a relevant avenue for future testing. Incipient evidence is present regarding the existence of periaxoplasmic ribosomal plaques, mRNA, and protein translation machinery in the axons, thus likely eliminating the bottleneck necessity of synaptic and axonal protein transport (55–62). Such evidence is virtually unknown for myenteric nitrergic neurons.

Though our report is novel from the perspective of being the first report to demonstrate significant reduction of myosin Va within varicosities in the peripheral nervous system, it has been earlier reported that myosin Va is significantly reduced in neuronal cell bodies in multiple brain regions several days after STZ-induced diabetes (63). One intriguing question that posits

from these observations is that if myosin Va is an important regulator of both vesicular and non-vesicular neurotransmission in both the central and peripheral nervous system, how can neuronal physiological function continue to operate, albeit at a lower level of performance, despite reduction, or absence of myosin Va. One likely reason of how neurotransmission function is sustained in the face of diminished myosin Va may be due to neuroplastic changes. For example, it has been reported that in the brain, similar diabetic conditions that result in myosin Va reduction also results in increased expression of non-muscle myosin II (NMMII) (64). NMMII plays a critical role in vesicular neurotransmitter release at the cortex of the nerve terminal. Whether NMMII plays important role in nitrergic neurotransmission merits to be investigated, though increasing argument is being placed for critical role of the cortical cytoskeleton in nitrergic neurotransmission (14). Proteomic analyses in whole gut tissue and protein identification in varicosities have shown the presence of heavy chain of NMMII (65), identified in the synaptosomes as Mg-dependent ATPase (66). Myosin light chain kinase (MLCK), the enzyme that activates NMMII, has been shown to be diminished in diabetes and this dysregulation reversed by insulin (67). However, the localization study was made in whole gut extracts and it is not known whether MLCK was down regulated in nerve terminals or smooth muscles or both.

Myosin Va expression in myenteric ganglia is much reduced in diabetic tissues in comparison to nNOS α and it seems likely that this selective reduction has taken place early on after diabetes induction. Earlier studies have actually shown increased nNOS mRNA in male diabetic tissues (9). It is possible that the transcription factor or factors that drive myosin Va genomic synthesis are exquisitely sensitive to the pathophysiological alterations resulting from STZ-induced diabetes. For example, down regulation of the snail transcription factor that binds to an E-box in the promoter region of myosin Va gene (68) may have inhibited myosin Va synthesis. It is increasingly recognized that persistent hyperglycemia can alter transcription because basal transcription operators and repressors are sensitive to intracellular glucose concentrations (36, 69, 70).

Independent reports have confirmed increased VIP-IR of nerve varicosities several weeks after STZ-induced diabetes (7, 71, 72), though studies in human diabetic tissues have not always confirmed these findings (38). While this may be related to neuroplastic changes resulting from decrease in nitrergic synthesis, the increased VIP-IR may also be related to diminution of vesicular exocytosis at the junctions (73), probably resulting from deficiency of the motor protein myosin Va. ATP exocytosis has been reported to be enhanced in diabetic enteric nerve terminals (74). This apparently confounding finding may be possible due to compensatory changes in purinergic signaling resulting from diminution in pre-junctional NO synthesis. Our observations of increased perikaryal volumes of some myosin Va neurons in diabetes may have resulted from attenuation of axonal transport. The relationship of stasis as a fundamental pathophysiology of diabetic neuropathy has long been recognized (75, 76).

Controlling hyperglycemia in the periphery through ethnopharmacologic glucosidase inhibitor obtained from *Pouteria* has the potential to reverse the loss of myosin Va in central nervous system

neurons (77). This is similar to previous observations of reversal of nitrergic neuropathy by insulin treatment (8, 32). Thus, there is a potential for pharmacological reversal of myosin Va transcriptional inhibition, at least in the early phases after STZ induction of diabetes.

In summary, the results of the present study show the dissociation between presences of the key inhibitory neurotransmitter synthesizing enzyme nNOS α in the nerve terminals but absence of its local transporter myosin Va in the jejunum of rats after 16 weeks of diabetes induction with STZ. Earlier studies with dilute mice have shown that nNOS α may be present in the nerve varicosities despite deficient myosin Va (12). Surprisingly, similar observations were made in diabetic nerve terminals in the present study. STZ is uptaken by cells with the aid of GLUT2 transporter (78). STZ causes promoter methylation, gene inactivation, and cellular toxicity (78). It may be possible that alterations of myosin Va expression may have resulted from STZ toxicity of myenteric ganglia. However, this possibility is low because we did not observe global inhibition of transcription or generalized cell loss. Low dose STZ was used for the current study in accordance with previous studies, which was used as model for diabetes induction (79–82). The validity of our observations may be confirmed from different models of diabetes including genetic and lipotoxic models, examining different regions of the gastrointestinal tract and the temporal progression of disease. This significant observation of deficient myosin Va in nerve terminals may potentially explain impairment of pre-junctional NO synthesis during EFS of diabetic gut neuromuscular strips despite presence of the nitrergic synthetic enzyme nNOS.

AUTHOR CONTRIBUTIONS

Arun Chaudhury, conceptualized project, image analyses, drafted manuscript; Marcilio Hubner De Miranda-Neto, supervised animal experiments; Renata Virginia Fernandes Pereira, performed animal experiments; Jacqueline Nelisis Zanoni, supervised animal experiments, image acquisition and project supervision; all authors read and approved final version of the manuscript.

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REFERENCES

- Xue L, Suzuki H. Electrical responses of gastric smooth muscles in streptozotocin-induced diabetic rats. *Am J Physiol* (1997) **272**:G77–83.
- Imaeda K, Takano H, Koshita M, Yamamoto Y, Joh T, Suzuki H. Electrical properties of colonic smooth muscle in spontaneously non-insulin-dependent diabetic rats. *J Smooth Muscle Res* (1998) **34**:1–11. doi:10.1540/jsmr.34.1
- Cellek S, Foxwell NA, Moncada S. Two phases of nitrergic neuropathy in streptozotocin-induced diabetic rats. *Diabetes* (2003) **52**:2353–62. doi:10.2337/diabetes.52.9.2353
- Martinez-Cuesta MA, Massuda H, Whittle BJ, Moncada S. Impairment of nitrergic-mediated relaxation of rat isolated duodenum by experimental diabetes. *Br J Pharmacol* (1995) **114**:919–24. doi:10.1111/j.1476-5381.1995.tb13291.x

5. Jenkinson KM, Reid JJ. Effect of diabetes on relaxations to non-adrenergic, non-cholinergic nerve stimulation in longitudinal muscle of the rat gastric fundus. *Br J Pharmacol* (1995) **116**:1551–6. doi:10.1111/j.1476-5381.1995.tb16372.x
6. Utkan T, Yildiz F, Utkan NZ, Gacar N, Göçmez SS, Ulak G, et al. Effects of diabetes and elevated glucose on nitrergic relaxations in the isolated duodenum of the rat. *Acta Diabetol* (2009) **46**:295–301. doi:10.1007/s00592-008-0086-z
7. Domènech A, Pasquinelli G, De Giorgio R, Gori A, Bosch F, Pumarola M, et al. Morphofunctional changes underlying intestinal dysmotility in diabetic RIP-1/hNF β transgenic mice. *Int J Exp Pathol* (2011) **92**:400–12. doi:10.1111/j.1365-2613.2011.00789.x
8. Watkins CC, Sawa A, Jaffrey S, Blackshaw S, Barrow RK, Snyder SH, et al. Insulin restores neuronal nitric oxide synthase expression and function that is lost in diabetic gastropathy. *J Clin Invest* (2000) **106**:373–84. doi:10.1172/JCI8273C1
9. Gangula PR, Maner WL, Micci MA, Garfield RE, Pasricha PJ. Diabetes induces sex-dependent changes in neuronal nitric oxide synthase dimerization and function in the rat gastric antrum. *Am J Physiol Gastrointest Liver Physiol* (2007) **292**:G725–33. doi:10.1152/ajpgi.00406.2006
10. Adegate E, al-Ramadi B, Saleh AM, Vijayarathay C, Ponery AS, Arafat K, et al. Increase in neuronal nitric oxide synthase content of the gastroduodenal tract of diabetic rats. *Cell Mol Life Sci* (2003) **60**:1172–9. doi:10.1007/s00018-003-2298-2
11. Way KJ, Young HM, Reid JJ. Diabetes does not alter the activity and localisation of nitric oxide synthase in the rat anococcygeus muscle. *J Auton Nerv Syst* (1999) **76**:35–44. doi:10.1016/S0165-1838(99)00005-3
12. Chaudhury A, He XD, Goyal RK. Myosin Va plays a key role in nitrergic neurotransmission by transporting nNOS α to enteric varicosity membrane. *Am J Physiol Gastrointest Liver Physiol* (2011) **301**:G498–507. doi:10.1152/ajpgi.00164.2011
13. Chaudhury A, Cristofaro V, Carew JA, Goyal RK, Sullivan MP. Myosin Va plays a role in nitrergic smooth muscle relaxation in gastric fundus and corpora cavernosa of penis. *PLoS One* (2014) **9**:e86778. doi:10.1371/journal.pone.0086778
14. Chaudhury A. Molecular handoffs in nitrergic neurotransmission. *Front Med* (2014) **1**:8. doi:10.3389/fmed.2014.00008
15. Chaudhury A, He XD, Goyal RK. Role of PSD95 in membrane association and catalytic activity of nNOS α in nitrergic varicosities in mice gut. *Am J Physiol Gastrointest Liver Physiol* (2009) **297**:G806–13. doi:10.1152/ajpgi.00279.2009
16. Pereira RV, de Miranda-Neto MH, da Silva Souza ID, Zanoni JN. Vitamin E supplementation in rats with experimental diabetes mellitus: analysis of myosin-V and nNOS immunoreactive myenteric neurons from terminal ileum. *J Mol Histol* (2008) **39**:595–603. doi:10.1007/s10735-008-9200-3
17. Espreafico EM, Cheney RE, Matteoli M, Nascimento AA, De Camilli PV, Larson RE, et al. Primary structure and cellular localization of chicken brain myosin-V (p190), an unconventional myosin with calmodulin light chains. *J Cell Biol* (1992) **119**:1541–57. doi:10.1083/jcb.119.6.1541
18. Wen Q, Chklovskii DB. A cost-benefit analysis of neuronal morphology. *J Neurophysiol* (2008) **99**:2320–8. doi:10.1152/jn.00280.2007
19. Cuntz H, Forstner F, Borst A, Häusser M. One rule to grow them all: a general theory of neuronal branching and its practical application. *PLoS Comput Biol* (2010) **6**(8):e1000877. doi:10.1371/journal.pcbi.1000877
20. Snider J, Pillai A, Stevens CF. A universal property of axonal and dendritic arbors. *Neuron* (2010) **66**:45–56. doi:10.1016/j.neuron.2010.02.013
21. Meijering E, Jacob M, Sarría JC, Steiner P, Hirling H, Unser M. Design and validation of a tool for neurite tracing and analysis in fluorescence microscopy images. *Cytometry A* (2004) **58**:167–76. doi:10.1002/cyto.a.20022
22. Hernandez L, Bazotte RB, Gama P, Miranda-Neto MH. Streptozotocin-induced diabetes duration is important to determine changes in the number and basophilicity of myenteric neurons. *Arq Neuropsiquiatr* (2000) **58**:1035–9. doi:10.1590/S0004-282X200000600010
23. Ramamurthy B, Höök P, Jones AD, Larsson L. Changes in myosin structure and function in response to glycation. *FASEB J* (2001) **15**:2415–22. doi:10.1096/fj.01-0183com
24. Katayama S, Haga Y, Saeki H. Loss of filament-forming ability of myosin by non-enzymatic glycosylation and its molecular mechanism. *FEBS Lett* (2004) **575**:9–13. doi:10.1016/j.febslet.2004.08.029
25. Szyrový I, Hodný Z. Non-enzymatic glycosylation of myosin: effects of diabetes and ageing. *Gen Physiol Biophys* (1992) **11**:301–7.
26. Brenman JE, Chao DS, Xia H, Aldape K, Brecht DS. Nitric oxide synthase complexed with dystrophin and absent from skeletal muscle sarcolemma in Duchenne muscular dystrophy. *Cell* (1995) **82**:743–52. doi:10.1016/0092-8674(95)90471-9
27. Macioce P, Gandolfi N, Leung CL, Chin SS, Malchiodi-Albedi F, Ceccarini M, et al. Characterization of NF-L and betaIIISigma1-spectrin interaction in live cells. *Exp Cell Res* (1999) **250**:142–54. doi:10.1006/excr.1999.4479
28. Alami NH, Jung P, Brown A. Myosin Va increases the efficiency of neurofilament transport by decreasing the duration of long-term pauses. *J Neurosci* (2009) **29**:6625–34. doi:10.1523/JNEUROSCI.3829-08.2009
29. Celikbilek A, Tanik N, Sabah S, Borekci E, Akylou L, Ak H, et al. Elevated neurofilament light chain (NFL) mRNA levels in prediabetic peripheral neuropathy. *Mol Biol Rep* (2014) **41**(6):4017–22. doi:10.1007/s11033-014-3270-y
30. Medori R, Autilio-Gambetti L, Monaco S, Gambetti P. Experimental diabetic neuropathy: impairment of slow transport with changes in axon cross-sectional area. *Proc Natl Acad Sci U S A* (1985) **82**:7716–20. doi:10.1073/pnas.82.22.7716
31. Stenkamp-Strahm CM, Kappmeyer AJ, Schmalz JT, Gericke M, Balemba O. High-fat diet ingestion correlates with neuropathy in the duodenum myenteric plexus of obese mice with symptoms of type 2 diabetes. *Cell Tissue Res* (2013) **354**:381–94. doi:10.1007/s00441-013-1681-z
32. Burnstock G, Mirsky R, Belai A. Reversal of nerve damage in streptozotocin-diabetic rats by acute application of insulin in vitro. *Clin Sci (Lond)* (1988) **75**:629–35.
33. Faussone-Pellegrini MS, Grover M, Pasricha PJ, Bernard CE, Lurken MS, Smyrk TC, et al. Ultrastructural differences between diabetic and idiopathic gastroparesis. *J Cell Mol Med* (2012) **16**:1573–81. doi:10.1111/j.1582-4934.2011.01451.x
34. Demedts I, Masaoka T, Kindt S, De Hertogh G, Geboes K, Farré R, et al. Gastrointestinal motility changes and myenteric plexus alterations in spontaneously diabetic biobreeding rats. *J Neurogastroenterol Motil* (2013) **19**:161–70. doi:10.5056/jnm.2013.19.2.161
35. Tronchini EA, Trevizan AR, Tashima CM, Pereira RV, Zanoni JN. Supplementation with 0.1% and 2% vitamin E in diabetic rats: analysis of myenteric neurons immunostained for myosin-V and nNOS in the jejunum. *Arq Gastroenterol* (2012) **49**:284–90. doi:10.1590/S0004-28032012000400010
36. Anitha M, Gondha C, Sutliff R, Parsadanian A, Mwangi S, Sitarman SV, et al. GDNF rescues hyperglycemia-induced diabetic enteric neuropathy through activation of the PI3K/Akt pathway. *J Clin Invest* (2006) **116**:344–56. doi:10.1172/JCI26295
37. Forrest A, Huizinga JD, Wang XY, Liu LW, Parsons M. Increase in stretch-induced rhythmic motor activity in the diabetic rat colon is associated with loss of ICC of the submuscular plexus. *Am J Physiol Gastrointest Liver Physiol* (2008) **294**:G315–26. doi:10.1152/ajpgi.00196.2007
38. Grover M, Farrugia G, Lurken MS, Bernard CE, Faussone-Pellegrini MS, Smyrk TC, et al. Cellular changes in diabetic and idiopathic gastroparesis. *Gastroenterology* (2011) **140**:1575–85.e8. doi:10.1053/j.gastro.2011.01.046
39. He CL, Soffer EE, Ferris CD, Walsh RM, Szurszewski JH, Farrugia G. Loss of interstitial cells of Cajal and inhibitory innervation in insulin-dependent diabetes. *Gastroenterology* (2001) **121**:427–34. doi:10.1053/gast.2001.26264
40. Gangula PR, Mukhopadhyay S, Pasricha PJ, Ravella K. Sepiapterin reverses the changes in gastric nNOS dimerization and function in diabetic gastroparesis. *Neurogastroenterol Motil* (2010) **22**(1325–31):e351–2. doi:10.1111/j.1365-2982.2010.01588.x
41. Heine CL, Kolesnik B, Schmidt R, Werner ER, Mayer B, Gorren AC. Interaction between neuronal nitric-oxide synthase and tetrahydrobiopterin revisited: studies on the nature and mechanism of tight pterin binding. *Biochemistry* (2014) **53**:1284–95. doi:10.1021/bi401307r
42. Huang JD, Brady ST, Richards BW, Stenolen D, Resau JH, Copeland NG, et al. Direct interaction of microtubule- and actin-based transport motors. *Nature* (1999) **397**:267–70. doi:10.1038/16722
43. Rao MV, Engle LJ, Mohan PS, Yuan A, Qiu D, Cataldo A, et al. Myosin Va binding to neurofilaments is essential for correct myosin Va distribution and transport and neurofilament density. *J Cell Biol* (2002) **159**:279–90. doi:10.1083/jcb.200205062
44. Chaudhury A, Rao YM, Goyal RK. PIN/LC8 is associated with cytosolic but not membrane-bound nNOS in the nitrergic varicosities of mice gut: implications for nitrergic neurotransmission. *Am J Physiol Gastrointest Liver Physiol* (2008) **295**:G442–51. doi:10.1152/ajpgi.90280.2008
45. Droz B, Koenig HL, Di Giamberardino L. Axonal migration of protein and glycoprotein to nerve endings. I. Radioautographic analysis of the renewal of protein in nerve endings of chicken ciliary ganglion after intracerebral injection

- of (3H)lysine. *Brain Res* (1973) **60**:93–127. doi:10.1016/0006-8993(73)90852-4
46. Droz B, Leblond CP. Axonal migration of proteins in the central nervous system and peripheral nerves as shown by radioautography. *J Comp Neurol* (1963) **121**:325–46. doi:10.1002/cne.901210304
 47. Hoffmann PN, Lasek RJ. The slow component of axonal transport. Identification of major structural polypeptides of the axon and their generality among mammalian neurons. *J Cell Biol* (1975) **66**:351–66. doi:10.1083/jcb.66.2.351
 48. Court FA, Álvarez J. Slow axoplasmic transport under scrutiny. *Biol Res* (2011) **44**:311–21. doi:10.4067/S0716-97602011000400001
 49. Álvarez J, Torres JC. Slow axoplasmic transport: a fiction? *J Theor Biol* (1985) **112**:627–51. doi:10.1016/S0022-5193(85)80028-X
 50. Miller KE, Heidemann SR. What is slow axonal transport? *Exp Cell Res* (2008) **314**:1981–90. doi:10.1016/j.yexcr.2008.03.004
 51. Roy S. Seeing the unseen: the hidden world of slow axonal transport. *Neuroscientist* (2014) **20**:71–81. doi:10.1177/1073858413498306
 52. Tang Y, Scott D, Das U, Gitler D, Ganguly A, Roy S. Fast vesicle transport is required for the slow axonal transport of synapsin. *J Neurosci* (2013) **33**:15362–75. doi:10.1523/JNEUROSCI.1148-13.2013
 53. Hirokawa N, Takemura R. Molecular motors and mechanisms of directional transport in neurons. *Nat Rev Neurosci* (2005) **6**:201–14. doi:10.1038/nrn1624
 54. Zhou R, Niwa S, Homma N, Takei Y, Hirokawa N. KIF26A is an unconventional kinesin and regulates GDNF-Ret signaling in enteric neuronal development. *Cell* (2009) **139**:802–13. doi:10.1016/j.cell.2009.10.023
 55. Calliari A, Sotelo-Silveira J, Costa MC, Nogueira J, Cameron LC, Kun A, et al. Myosin Va is locally synthesized following nerve injury. *Cell Motil Cytoskeleton* (2002) **51**:169–76. doi:10.1002/cm.10017
 56. Campenot RB, Eng H. Protein synthesis in axons and its possible functions. *J Neurocytol* (2000) **29**:793–8. doi:10.1023/A:1010939307434
 57. Edstrom A, Edstrom JE, Hokfelt T. Sedimentation analysis of ribonucleic acid extracted from isolated Mauthner nerve fibre components. *J Neurochem* (1969) **16**:53–66. doi:10.1111/j.1471-4159.1969.tb10343.x
 58. Gumy LE, Tan CL, Fawcett JW. The role of local protein synthesis and degradation in axon regeneration. *Exp Neurol* (2010) **223**:28–37. doi:10.1016/j.expneurol.2009.06.004
 59. Kun A, Otero L, Sotelo-Silveira JR, Sotelo JR. Ribosomal distributions in axons of mammalian myelinated fibers. *J Neurosci Res* (2007) **85**:2087–98. doi:10.1002/jnr.21340
 60. Toth CC, Willis D, Twiss JL, Walsh S, Martinez JA, Liu W-Q, et al. Locally synthesized calcitonin gene related peptide has a critical role in peripheral nerve regeneration. *J Neuropathol Exp Neurol* (2009) **68**:326–37. doi:10.1097/NEN.0b013e31819ac71b
 61. Weiss PA. Neuronal dynamics and axonal flow. V. The semisolid state of the moving axonal column. *Proc Natl Acad Sci U S A* (1972) **69**:620–3. doi:10.1073/pnas.69.5.1309
 62. Willard M, Cowan WM, Vagelos PR. The polypeptide composition of intraxonally transported proteins: evidence for four transport velocities. *Proc Natl Acad Sci U S A* (1974) **71**:2183–7. doi:10.1073/pnas.71.6.2183
 63. da Costa AV, Calábria LK, Nascimento R, Carvalho WJ, Goulart LR, Espindola FS. The streptozotocin-induced rat model of diabetes mellitus evidences significant reduction of myosin-Va expression in the brain. *Metab Brain Dis* (2011) **26**:247–51. doi:10.1007/s11011-011-9259-5
 64. Calábria LK, da Cruz GC, Nascimento R, Carvalho WJ, de Gouveia NM, Alves FV, et al. Overexpression of myosin-IIb in the brain of a rat model of streptozotocin-induced diabetes. *J Neurol Sci* (2011) **303**:43–9. doi:10.1016/j.jns.2011.01.017.65
 65. Marvin-Guy L, Lopes LV, Affolter M, Courtet-Compondu MC, Wagnière S, Bergonzelli GE, et al. Proteomics of the rat gut: analysis of the myenteric plexus-longitudinal muscle preparation. *Proteomics* (2005) **5**:2561–9. doi:10.1002/pmic.200401265
 66. Agoston DV, Whittaker VP. Characterization, by size, density, osmotic fragility, and immunoaffinity, of acetylcholine- and vasoactive intestinal polypeptide-containing storage particles from myenteric neurones of the guinea-pig. *J Neurochem* (1989) **52**(5):1474–80. doi:10.1111/j.1471-4159.1989.tb09196.x
 67. Hu W, Feng P. Myosin light chain kinase is involved in the mechanism of gastrointestinal dysfunction in diabetic rats. *Dig Dis Sci* (2012) **57**:1197–202. doi:10.1007/s10620-012-2041-7
 68. Lan L, Han H, Zuo H, Chen Z, Du Y, Zhao W, et al. Upregulation of myosin Va by snail is involved in cancer cell migration and metastasis. *Int J Cancer* (2010) **126**:53–64. doi:10.1002/ijc.24641
 69. Issad T, Kuo M. O-GlcNAc modification of transcription factors, glucose sensing and glucotoxicity. *Trends Endocrinol Metab* (2008) **19**:380–9. doi:10.1016/j.tem.2008.09.001
 70. Kamemura K, Hart GW. Dynamic interplay between O-glycosylation and O-phosphorylation of nucleocytoplasmic proteins: a new paradigm for metabolic control of signal transduction and transcription. *Prog Nucl Acid Res Mol Biol* (2003) **73**:107–36. doi:10.1016/S0079-6603(03)01004-3
 71. Belai A, Lincoln J, Milner P, Crowe R, Loesch A, Burnstock G. Enteric nerves in diabetic rats: increase in vasoactive intestinal polypeptide but not substance P. *Gastroenterology* (1985) **89**:967–76.
 72. Hermes-Uliana C, Panizzon CP, Trevizan AR, Sehaber CC, Ramalho FV, Martins HA, et al. Is L-glutathione more effective than L-glutamine in preventing enteric diabetic neuropathy? *Dig Dis Sci* (2014) **59**:937–48. doi:10.1007/s10620-013-2993-2
 73. Belai A, Lincoln J, Burnstock G. Lack of release of vasoactive intestinal polypeptide and calcitonin gene-related peptide during electrical stimulation of enteric nerves in streptozotocin-diabetic rats. *Gastroenterology* (1987) **93**:1034–40.
 74. Belai A, Lefebvre RA, Burnstock G. Motor activity and neurotransmitter release in the gastric fundus of streptozotocin-diabetic rats. *Eur J Pharmacol* (1991) **194**:225–34. doi:10.1016/0014-2999(91)90109-4
 75. Jakobsen J. Axonal dwindling in early experimental diabetes. I. A study of cross sectioned nerves. *Diabetologia* (1976) **12**:539–46. doi:10.1007/BF01220629
 76. Carpenter S. Proximal axonal enlargement in motor neuron disease. *Neurology* (1968) **18**:841–51. doi:10.1212/WNL.18.9.841
 77. da Costa AV, Calábria LK, Furtado FB, de Gouveia NM, Oliveira RJ, de Oliveira VN, et al. Neuroprotective effects of *Pouteria ramiflora* (Mart.) Radlk (Sapotaceae) extract on the brains of rats with streptozotocin-induced diabetes. *Metab Brain Dis* (2013) **28**:411–9. doi:10.1007/s11011-013-9390-6
 78. Ventura-Sobrevilla J, Boone-Villa VD, Aguilar CN, Román-Ramos R, Vega-Avila E, Campos-Sepúlveda E, et al. Effect of varying dose and administration of streptozotocin on blood sugar in male CD1 mice. *Proc West Pharmacol Soc* (2011) **54**:5–9.
 79. Like AA, Rossini AA. Streptozotocin-induced pancreatic insulinitis: new model of diabetes mellitus. *Science* (1976) **193**:415–7. doi:10.1126/science.180605
 80. Wilson GL, Leiter EH. Streptozotocin interactions with pancreatic beta cells and the induction of insulin-dependent diabetes. *Curr Top Microbiol Immunol* (1990) **156**:27–54.
 81. Sullivan KA, Hayes JM, Wiggin TD, Backus C, Su Oh S, Lentz SI, et al. Mouse models of diabetic neuropathy. *Neurobiol Dis* (2007) **28**:276–85. doi:10.1016/j.nbd.2007.07.022
 82. Tesch GH, Allen TJ. Rodent models of streptozotocin-induced diabetic nephropathy. *Nephrology (Carlton)* (2007) **12**:261–6. doi:10.1111/j.1440-1797.2007.00796.x

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