Axotomy-Induced Changes in the Chemical Coding Pattern of Colon Projecting Calbindin-Positive Neurons in the Inferior Mesenteric Ganglia of the Pig

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Abstract The present study examines the response of colon-projecting neurons localized in the inferior mesenteric ganglia (IMG) to axotomy in the pig animal model. In all animals (n=8), a median laparotomy was performed under anesthesia and the retrograde tracer Fast Blue was injected into the descending colon wall. In experimental animals (n=4), the descending colon was exposed and the bilateral caudal colonic nerves were identified and severed. All animals were euthanized and the inferior mesenteric ganglia were harvested and processed for double-labeling immunofluorescence for calbindin-D28k (CB) in combination with either tyrosine hydroxylase (TH), neuropeptide Y (NPY), somatostatin (SOM), vasoactive intestinal polypeptide (VIP), nitric oxide synthase (NOS), Leu-enkephalin (LENK), substance P, vesicular acetylcholine transporter, or galanin. Immunohistochemistry revealed significant changes in the chemical coding pattern of injured inferior mesenteric ganglion neurons. In control animals, Fast Blue-positive neurons were immunoreactive to TH, NPY, SOM, VIP, NOS, LENK, and CB. In the experimental group, the numbers of TH-, NPY-, and SOM-expressing neurons were reduced, whereas the number of neurons immunoreactive to LENK was increased. Our data indicate that the colon-projecting neurons of the porcine IMG react to the axotomy in a similar, but not an identical manner in a comparison to other species, especially rodents. Further studies are needed to elucidate the detailed factors/mechanisms involved in the response to nerve injury.

Keywords Axotomy · Neuropeptides · Colon-supplying neurons · Inferior mesenteric ganglion · Pig

Introduction

The blood flow, secretions, and motility of the gastrointestinal tract (GIT) are controlled by the enteric nervous system. It is composed of both the intramural plexuses and extrinsic autonomic neurons that are located in the prevertebral and paravertebral ganglia (Brown and
Timmermans 2004; Furness 2012; Luckensmeyer and Keast 1995). Previous investigations have shown that in the case of the colon, the main source of extrinsic innervation is located within inferior mesenteric ganglion (IMG) (Dalsgaard and Elfvin 1982; Li and Masuko 2001). Moreover, both enteric neurons and extrinsic innervation of the gastrointestinal tract are able to undergo structural, functional, and chemical changes as a result of adaptive or reparative processes in response to different physiological and pathological stimuli, such as development, aging, diet, or intestinal and extra-intestinal diseases (Balemba et al. 1998; Ekblad et al. 1998; Phillips and Powley 2007).

One of the factors that induces a sequence of morphological, biochemical, and electrophysiological changes both in affected neurons and neighboring nonneuronal cells is axotomy (Sun and Zigmond 1996; Yamada et al. 2008). It has been shown that the transection of the axon leads to a decrease of catecholamine biosynthesis and changes in the expression pattern of neuroprotective/neuroregenerative peptides. These include galanin (GAL), nitric oxide synthase (NOS), vasoactive intestinal peptide (VIP), and substance P (SP) (Hyatt-Sachs et al. 1996; Klimaschewski et al. 1996; Reimer and Kanje 1999; Zigmond and Sun 1997; Zigmond 2000). These changes are an example of plasticity in the peripheral nervous system that allows the sympathetic neurons to survive and regenerate after axonal damage (Boeshore et al. 2004; Sun et al. 1996; Zigmond and Sun 1997).

Although a decrease of catecholamine biosynthesis is commonly observed after injury to sympathetic neurons (Skobowiat et al. 2011; Zigmond 2000), the second phenomenon, i.e., the induction or overexpression of neuroprotective/neuroregenerative peptides in affected neurons seems to be organ dependent as well as species specific (Lindh et al. 1993; Majewski et al. 2002; Olsson et al. 2006; Reimer and Kanje 1999; Skobowiat et al. 2011; Zigmond 1997). For example, in the rat, NOS and VIP expression rates increase in a subpopulation of neurons of the axotomized superior cervical ganglion (Hyatt-Sachs et al. 1996; Klimaschewski et al. 1994, 1996; Mohney et al. 1994), but in the pig, axotomy did not change NOS or VIP synthesis rate in neurons of the porcine inferior mesenteric ganglion (Majewski et al. 2002). On the other hand, axotomy changes the chemical coding of neuronal cells within the enteric nervous system in the descending colon (Gonkowski and Calka 2010, 2012).

Apart from the decrease of catecholamine biosynthesis and the changes in the expression pattern of neuroprotective/neuroregenerative peptides, axotomy also causes a massive influx of calcium into the lesioned neurons (Obal et al. 2006; Wolf et al. 2001). Calcium influx is highly cytotoxic, so resection of the nerve should induce extensive neuronal cell death (LoPachin and Lehning 1997). However, axotomy of the hypoglossal nerve does not induce extensive neuronal cell death in rats (Krebs et al. 1997). A large percentage (70%) of these neurons has been shown to survive despite permanent target deprivation (Krebs et al. 1997). One of the mechanisms responsible for this substantial survival rate in rats may be changes in the Ca\(^2+\) buffering system. For example, as a reaction to the hypoglossal nerve axotomy, lesioned neurons trigger an increase in calbindin expression which acts as a compensatory Ca\(^2+\) buffering system, enabling neurons to maintain Ca\(^2+\) homeostasis and the survival after axotomy (Krebs et al. 1997). Calbindin-D28K is an intracellular calcium-binding protein, which acts as Ca\(^2+\) buffering system in the cytoplasm (Schwaller et al. 2002; Schwaller 2009). By means of this property, calbindin may protect neurons against large fluctuations in free intracellular Ca\(^2+\) and may prevent cell death. Therefore, we designed our experiments to decipher changes induced by caudal colonic nerve axotomy on the chemical coding of postganglionic, sympathetic IMG neurons innervating the colon of the pig by applying Fast Blue tracing, axotomy, and double-labeling immunofluorescence for calbindin-D28k in combination with various other substances. It should be pointed out that axotomy of the neurons supplying the GIT can be associated with the majority of surgical operations on stomach and intestines, but also with the mechanical injuries of abdominal cavity. So the studies on the axotomy-induced changes in the chemical coding of neurons supplying the gastrointestinal tract seem to be of interest, especially in the pig due to its anatomical, histological, and physiological similarities to the human GI tract (Brown and Timmermans 2004). Pigs and humans also have similar immune systems and inflammatory responses (Patterson et al. 2008; Swindle and Smith 2007; Verma et al. 2011). Due to all these similarities, the pig animal model is an optimal model for various human diseases including cardiovascular diseases, obesity, diabetes, as well as injury and repair of tissues (Litten-Brown et al. 2010; Sullivan et al. 2001; Swindle and Smith 2007; Swindle et al. 2012). Therefore, as calbindin-D28k (CB) could play an important role as a calcium ion buffer, especially under pathological conditions, we decided to investigate in detail the eventual changes in the number and chemical phenotypes of colon-projecting and CB-expressing neurons in IMG induced by an axotomy.

Material and Methods

Study Subjects

The study was performed on eight immature female pigs of the Large White Polish breed (8 weeks old) divided into two groups: control (C group, \(n=4\)) and axotomized animals (AXO group; \(n=4\)). All animals were housed and treated in accordance with rules approved by the ethics committee.
Anesthesia and Surgery

Surgery was performed under fractionated thiobarbital (Thiopental, Sandoz, Austria; 20 mg/kg b.w., i.v.) anesthesia. Prior to administration (30 min) of the main anesthetic, the animals were pretreated with atropine sulfate (Polfa, Poland; 0.04 mg/kg b.w., s.c.) and azaperone (Stresnil, Janssen Pharmaceutica, Belgium; 2.0 mg/kg b.w., i.m.). All animals (n=8) were laparotomized and injected with 5 % aqueous solution of the fluorescence retrograde neuronal tracer Fast Blue (FB; Dr. K. Illing GmbH, Gross-Umstadt, Germany) into the wall of the descending colon. A total volume of 40 μl of 5 % aqueous dye solution was injected into the exposed descending colon using a Hamilton syringe with a 26-gauge needle. Multiple injections (n=40; 1 μl each) were made into the right side of the descending colon wall, using the mesocolon as the bordering mark. Three weeks later, all animals underwent a median laparotomy to expose the descending colon and the bilateral nervi colici caudales (a sole link between the IMG and this bowel segment). In half of the animals, the nerves were transected (AXO group), while in the control group after visualization their continuity was maintained. After 7 days, all animals were re-anesthetized and euthanized by an overdose of sodium thiobarbital then perfused transcardially with 4 % buffered paraformaldehyde (pH 7.4). Following perfusion, small tissue blocks containing the IMG were collected from all studied animals and postfixed by immersion in the same fixative for 4 h, washed twice in 0.1 M phosphate buffer (pH=7.4, 4 °C), and then stored in 18 % sucrose at 4 °C until sectioning.

Immunofluorescence Experiments

Ten-micrometer-thick cryostat sections of IMGs were processed for routine double-immunofluorescence labeling using primary antisera raised in different species and species-specific secondary antibodies (Table 1). All sections were incubated overnight in a humid chamber with a mixture consisting of antibodies directed towards CB and an antiserum raised against either tyrosine hydroxylase (TH), neuropeptide Y (NPY), somatostatin (SOM), vasoactive intestinal polypeptide (VIP), nitric oxide synthase (NOS), leucine-enkephalin (LENK), SP, vesicular acetylcholine transporter (VACHT), or GAL (Table 1). Sections were then incubated with a mixture of appropriate fluorescein isothiocyanate (FITC)-conjugated secondary antisera and biotinylated goat antirabbit antibodies (1 h). The latter antibodies were finally visualized by additional incubation of sections with streptavidin–CY3 complex (1 h). After staining, the sections were mounted with carbonate-buffered glycerol (pH 8.6) and cover slipped. Each step of immunolabeling was followed by rinsing of the sections with PBS (3×15 min).

Controls

Standard controls, i.e., preabsorption for the neuropeptide antisera (20 μg of appropriate antigen per 1 ml of corresponding antibody at working dilution), and the omission and replacement of all primary antisera by nonimmune sera or PBS were applied to test both antibody and method specificity. Antigens used for the preabsorption tests were as follows: SOM (code S9129, Sigma, St. Louis, USA), NPY (code PEP-87135, Dianova, Hamburg, Germany), and SP (code S6883, Sigma). Apart from the preabsorption experiments, all primary antibodies were tested on selected porcine brain tissue samples where the studied antigens were previously reported. The secondary antisera were tested by omission and replacement procedures.

Cell Counting and Statistics

The sections were viewed under an Olympus BX51 fluorescence microscope equipped with a barrier filter for FB and the respective filters for FITC and CY3. Microphotographs were acquired with a CCD camera connected by a PC equipped with Olympus AnalySIS image analysis software (ver. 3.2; Soft Imaging System GmbH, Germany). To determine the relative number of FB-positive cells, the neurons were counted in every 16th section in the whole ganglion to avoid double counting (approximately ten sections per animal, per combination studied). The number of cells present on the particular sections as well as the relative frequency of cells exhibiting particular neurochemical phenotype was analyzed by the first author and two other co-authors, blinded to the identity of the antigens studied. At least 560 FB+ neurons were analyzed for any combination of antigens. Only neurons with clearly visible nucleus (seen as non-immunoreactive “hole” in the middle of the perikaryon) were counted. Data pooled from particular animals groups were expressed as means±standard deviation of mean and analyzed with chi-square tests. Statistical significance was inferred at
Results

The control animals contained a large number of retrogradely labeled neurons (FB+) that were distributed bilaterally, i.e., within both the left and right ganglia. The majority of FB+ cells were TH or NPY immunoreactive (87.8±4.9 and 84±3.1 %, respectively) and a large percentage of them co-expressed CB (36.5±3.9 and 28.9±3.4 %, respectively; Table 2 and Figs. 1 and 2a–h). In addition, small subpopulations of FB+ neurons co-expressed SOM (18.7±2.2 %), VIP (5±0.9 %), NOS (1.8±0.8 %), and LENK (1.4±0.8 %; Table 2 and Figs. 1, 2i–l, and 3a–l). However, none of the FB+ perikarya were found to be immunopositive to SP, VAChT, and GAL (Table 2). In axotomized animals, the total number of FB+ neurons did not change; however, the coding patterns of these cells were profoundly changed. Thus, a strong reduction was observed in the

| Antigen | Code | Host species | Dilution | Supplier |
|---------|------|--------------|----------|----------|
| Primary antibodies | | | | |
| CB | CB-38 | Rabbit | 1:20,000 | SWANT |
| TH | MAB 318 | Mouse | 1:80 | Chemicon |
| NPY | NZ1115 | Rat | 1:300 | Biomol |
| SOM | 8330-0009 | Rat | 1:100 | Biogenesis |
| VIP | 9535-0504 | Mouse | 1:2,000 | Biogenesis |
| NOS | N2280 | Mouse | 1:2,000 | Sigma |
| LENK | 4140-0355 | Mouse | 1:1,000 | Biogenesis |
| SP | 8450-0505 | Rat | 1:300 | Biogenesis |
| VAChT | H-V007 | Goat | 1:2,000 | Phoenix |
| GAL | T-5036 | Guinea pig | 1:1,000 | Peninsula |
| Secondary reagents | | | | |
| Donkey anti-mouse IgG (H+L) conjugated with FITC | | | 1:800 | Jackson |
| Donkey anti-rat IgG (H+L) conjugated with FITC | | | 1:800 | Jackson |
| Donkey anti-guinea pig IgG (H+L) conjugated with FITC | | | 1:1,000 | Jackson |
| Donkey anti-goat IgG (H+L) conjugated with FITC | | | 1:1,000 | Jackson |
| Biotinylated goat antirabbit immunoglobulins | | | 1:1,000 | DAKO |
| Biotin conjugated F(ab) fragment anti-rabbit IgG | | | 1:1,000 | BioTrend |
| CY3-conjugated streptavidin | | | 1:9,000 | Jackson |

Data were expressed as mean±standard deviation (SD)

*S studied substance (NPY, SOM, VIP, etc.)
number of FB+/TH+ (61.0±3.5 vs. 87.8±4.9 %, AXO vs. control, respectively; \( P \leq 0.001 \)), FB+/NPY+ (43±1.0 vs. 84±3.1 %, AXO vs. control, respectively; \( P \leq 0.001 \)), and FB+/SOM+ (3.7±0.2 vs. 18.7±2.2 %, AXO vs. control, respectively; \( P \leq 0.001 \)) neurons (Table 2 and Figs. 1 and 2a–l). Alternatively, the number of FB+/VIP+ (\( P > 0.05 \)), FB+/NOS+ (\( P > 0.05 \)), and FB+/CB+ (\( P > 0.05 \)) neurons was not changed, while the number of FB+/LENK+ (5.8±1.6 vs. 1.4±0.8 %, AXO vs. control, respectively; \( P \leq 0.001 \)) cells increased (Table 2 and Figs. 1 and 3a–l).

It has been found that under physiological conditions, FB+/CB+ neurons constituted approximately 35 % of the whole FB+ cellular population. It should be stressed that these cells were also neurochemically heterogeneous (Table 2 and Figs. 1 and 2). For example, many of FB+/CB+ cells co-expressed TH (33.4±4.7 %) either NPY (25.1±3.8 %) or SOM (11.1±1.3 %), while a few of them co-expressed VIP (1.5±1.0 %; Table 2 and Figs. 1, 2a–h, and 3a–d). None of the FB+/CB+ positive perikarya were found to be immunopositive to NOS, LENK, SP, VACHT, and GAL (Table 2 and Figs. 1 and 3e–l). In axotomized animals, although the total number of the FB+/CB+ cells did not change, a strong reduction was observed among FB+/CB+/TH+ (16.8±3.5 vs. 33.4±4.7 %, AXO vs. control, respectively; \( P \leq 0.001 \)), FB+/CB+/NPY+ (1.4±0.4 vs. 25.1±3.8 %, AXO vs. control, respectively; \( P \leq 0.001 \)), and FB+/CB+/SOM+ (3.4±
Fig. 2 Representative images of colon-projecting neurons located in IMG of the control animals (C, a–l) and axotomized animals (AXO, a’–l’). d, d’, h, h’, l, and l’ are composites of merged images taken separately from blue (a, a’, e, e’, i, i’), red (b, b’, f, f’, j, j’), and green (c, c’, g, g’, k, k’) fluorescent channels. Scale bar in each figure = 50 μm. a–d and a’–d’ FB+/CB+/TH+ neurons in C and AXO animals, respectively. The FB+ (a and a’) and FB+/CB+/TH+ (d and d’) neurons in C and AXO animals, respectively (small arrows). Some FB+/TH+ neurons in C group were CB negative (long arrow), FB+/CB+/TH+ neurons in AXO animals (small double arrows). e–h and e’–h’ FB+/CB+/NPY+ neurons in C and AXO animals, respectively. Note strong reduction number of FB+/CB+/NPY+ (h and h’) neurons in AXO animals (small arrow), FB+/CB+/NPY+ (small arrow), FB+/CB+/NPY+ (small double arrows), and FB+/CB+/NPY+ (long arrows). i–l and i’–l’ FB+/CB+/SOM+ neurons in C and AXO animals, respectively. Note a strong reduction in the number of SOM+ (k and k’) and FB+/CB+/SOM+ (l and l’) neurons in AXO animals (small arrows). FB+/CB+/SOM+ (small arrows), FB+/CB+/SOM+ (small double arrows).
Fig. 3 Representative images of colon-projecting neurons located in IMG of the control animals (C, a–l) and axotomized animals (AXO, a′–l′). d, d′, h, h′, l, and l′ are composites of merged images taken separately from blue (a, a′, e, e′, i, i′), red (b, b′, f, f′, j, j′), and green (c, c′, g, g′, k, k′) fluorescent channels. Scale bar in each figure=50 μm. a–d and a′–d′ FB+/CB+/VIP+ neurons in C and AXO animals, respectively. FB+/CB+/VIP+ (small arrow), FB+/CB+/VIP− (small double arrows). e–h and e′–h′ Note the lack of FB+/CB+/NOS+ (h and h′) neurons in C and AXO animals and strong reduction in the number of NOS+ (g and g′) neurons in AXO animals. FB+/CB+/NOS− (long arrow), FB+/CB−/NOS− (long double arrows). i–l and i′–l′ Note the lack of FB+/CB+/LENK+ (l and l′) neurons in C and AXO animals. FB+/CB−/LENK− (long double arrows), FB+/CB−/LENK+ (long arrows).
0.1 vs. 11.1±1.3 %, AXO vs. control, respectively; P≤0.05) neurons (Table 2 and Figs. 1 and 2a′–l).

Discussion

The pig pattern of axotomy-induced changes in the chemical coding of IMG neurons supplying descending colon The present study demonstrates changes in the chemical coding of the colon-projecting neurons located in the porcine IMG following axotomy of the nervi colici caudales. These changes include a reduction in the number of neurons expressing TH, NPY, and SOM and an increase in the number of neurons immunoreactive to LENK. Although the number of CB⁻ neurons was similar in both the control and axotomized animals, there were significant discrepancies concerning the neurochemical features of this neuronal sub-set prior and after the injury. Thus, we have observed a strong downregulation of TH, NPY, and SOM expression in FB⁺/CB⁻ neurons. Calbindin-D28K plays a major role in calcium homeostasis in neurons and other cell types acting as a fast Ca²⁺ buffering system in the cytoplasm (Schwaller et al. 2002; Schwaller 2009). This way, calbindin may protect neurons against large fluctuations in free intracellular Ca²⁺ and prevent cell death. Since axotomy causes a massive influx of calcium into the lesioned neurons (Wolf et al. 2001), an increase in calbindin expression in IMG should be expected. However, it seems not to be the case as the number of CB-expressing neurons was similar in both the control and axotomized animals. One of the possible explanations for such phenomenon in IMG may be that the other calcium-binding proteins like parvalbumin or calretinin were engaged. Such mechanism, for example, i.e., ability to upregulate parvalbumin after axotomy, paralleled by a smaller increase of intracellular calcium was reported in oculomotor neurons of mice (Obal et al. 2006).

The pig pattern of axotomy-induced changes in the IMG vs. other ganglia and/or species It is widely accepted that one of the most relevant changes in the neuronal phenotype following axotomy is the downregulation of physiological neurotransmitter production and the increase in the expression of neuropeptides which are essential for survival and/or regeneration (Hyatt-Sachs et al. 1996; Zigmund and Sun 1997; Zigmund 2000). Our data indicate that the colon-projecting neurons located in the porcine IMG react in a similar manner; however, this manner differs in some details from that described in other ganglia and/or species.

TH The considerable decrease in TH expression in the FB⁺ population in porcine IMG after caudal colonic nerve axotomy is well in line with earlier data obtained from the porcine IMG after partial or total uterus extirpation (Wasowicz 2003a, b, c). The same phenomenon was also observed in the rat superior cervical ganglia (SCG), where the decreased expression of the catecholamine-producing enzymes has also been noticed after axotomy (Klimaschewski et al. 1996; Shadiack et al. 2001; Sun and Zigmund 1996).

NPY In addition to the decreased catecholamine production, the axotomy-induced reduction in the number of NPY perikarya was also observed in the porcine IMG, which is consistent with the data obtained from the rat SCG described earlier by Bachoo et al. (1992) and Sun and Zigmund (1996). A decrease of NPY expression should not be surprising since NPY-expressing neurons form a large population among IMG neurons and many of them co-express TH (Pidsudko et al. 2008). Interestingly, after a partial or total uterus extirpation, uterus-projecting IMG neurons expressing NPY were upregulated (Wasowicz 2003a).

SOM SOM is another substance in the present study which was downregulated in neurons of the porcine IMG after axotomy. The reduction of SOM expression in FB⁺ perikarya was surprising since SOM modulates K⁺/Ca²⁺ channel activities that “sensitize” the sympathetic neurons to nerve growth factor and thus can facilitate the neuroregeneration (Patel 1999). Moreover, the induction of SOM production after axotomy has been described previously in the porcine sympathetic chain ganglia (Skobowiat et al. 2011).

VIP and NOS VIP and NOS expressions were not changed in neurons of the porcine IMG after colonic nerve axotomy which is consistent with the previous reports on this subject (Majewski et al. 2002). However, these data run counter to the results of the injury-related responses observed in the neurons of various sympathetic ganglia in rodents, suggesting the existence of either species or target tissue-dependent differences in injury-induced responses (Hyatt-Sachs et al. 1996; Klimaschewski et al. 1996; Shadiack et al. 2001; Sun and Zigmund 1996). For example, it is widely accepted that the axotomy induces or increases the number of neurons expressing VIP and/or NOS in sympathetic ganglia in rodents (Hyatt-Sachs et al. 1996; Klimaschewski et al. 1996; Shadiack et al. 2001; Sun and Zigmund 1996). This phenomenon may be attributable to both continuous release of leukemia inhibitory factor from nonneuronal cells at the site of injury (Zigmund 2000) and nerve growth factor deprivation of the affected cells (Shadiack et al. 2001). However, this mechanism does not appear to be involved in the axotomy-driven changes seen in the present study as we did not observed any increase in the number of VIP and/or NOS neurons in the porcine IMG.

LENK Among all IMG neurons studied in the present experiment, only these expressing LENK became more numerous after caudal colonic nerves axotomy. LENK
overexpression in response to axotomy was already reported in the IMG of the pigs (Majewski et al. 2002). However, axotomy is able to induce increases, but also decreases in the number of LENK expressing cells, depending on the model used (Back and Gorenstein 1994; Linda et al. 1990). Most probably, it is the target tissue that determines the regulation of enkephalins, i.e., target-derived factors either suppress or activate the expression of the leu-enkephalin mRNA precursor normally. Axotomy prevents the target contact, so the target-derived factors cannot act on gene expression anymore.

Summary Our data indicate that the colon-projecting neurons of the porcine IMG react to the axotomy in a similar but not identical manner in a comparison to other species, especially rodents. The existence of species and target tissue-dependent differences in injury-induced responses suggest that further studies are needed to elucidate the detailed factors/mechanisms involved in the response to nerve injury.

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