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We use this protocol and it's working

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Tache_Mulugeta_OT2OD024899_Thoracolumbar and sacral nerve roots acute electrical stimulation and colonic motility measurements V.1

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ABSTRACT

This protocol describes a process for the measurement of acute electrical stimulation-induced effects on colonic motility in anesthetized young adult Yucatan minipigs. Signals recorded from manometry probes inserted into the proximal, transverse and distal colonic regions were used to measure the effect of stimulation on the sacral or thoracolumbar nerve roots in an acute anesthetized preparation. The effect of stimulation was quantified as motility index assessments before, during and after stimulation, and the data was used to create a functional map of colonic motor response to spinal nerve roots stimulation.

MATERIALS

Mikro-CathTM Pressure Catheter, Millar Inc., Ref# 825-0101
https://millar.com/Clinical/MikroCath/#quicktabs-mikro_cath_diagnostic_pressure_c=2

PEC-10D Pressure cable, Millar Inc., Ref# 850-5090 <https://millar.com/Clinical/Control-Units-and-Accessories/>

PCU-2000, Dual Channel Pressure Control Unit, Millar Inc., Ref# 880-0129
<https://millar.com/Clinical/Control-Units-and-Accessories/>

Micro 1401, Data Acquisition Unit, Cambridge Electronic Design, Ltd, Ref# CED Micro1401
<http://ced.co.uk/products/mic4in>

Spike 2 7.10, Windows 7, Cambridge Electronic Design, Ltd, <http://ced.co.uk/products/spkovin>

Animals

- 1 Six-to-seven months old (25-36 kg) male Yucatan minipigs (S&S farms, Ramona, CA), castrated at 7 days of age, are group housed in pens (either bedding or grate floor depending on housing availabilities - 2 pigs/pen, 42ft²) in an environmentally controlled room (lights on/off 6AM/6PM, 61-81°F) under SPF conditions.

☒ Yucatan minipig **S&S Farms**

All pigs were offered ad libitum access to diet (5p94 Prolab mini pig diet, PMI nutrition) and filtered tap water.

All husbandry practices and procedures conformed to the NIH Guide for the Care and Use of Laboratory Animals (8th edition) and were reviewed and approved by the UCLA Animal Research Committee (Institutional Animal Care and Use Committee). All efforts were made to minimize any suffering as well as the number of animals used.

Surgical Procedures

- 2 Pigs were fasted for at least 12h prior to surgery with free access to water.

For surgical level anesthesia, pigs were premedicated with midazolam (1 mg/kg),

☒ Midazolam HCl Injection, 50mg/10mL (5mg/mL), C4 **Henry Schein Animal Health Catalog #067695**
ketamine (15 mg/kg)

☒ Ketamine HCl Injection, 200mg/20mL (10mg/mL), C3N **Henry Schein Animal Health Catalog #06831**
and meloxicam (0.3 mg/kg)

☒ EloxiJect (Meloxicam) Injection, 5mg/mL **Henry Schein Animal Health Catalog #049755** injected intramuscularly.

They were then intubated, connected to a respirator for ventilation (breathing rate maintained between 13-16 breaths/min), and maintained under general anesthesia with 1-3% inhaled isoflurane.

☒ IsoThesia (Isoflurane) Solution **Henry Schein Animal Health Catalog #029405**

Maintenance fluids (lactated ringers) were administered at 10 ml/kg/h.

☒ Lactated Ringers Injection, USP, Preservative-Free, Baxter **Henry Schein Animal Health Catalog #059380**

During the first part of the surgical procedure, pigs were positioned on a heating pad (32°C) in supine position.

A femoral heart line was placed. When needed, the heart line was flushed with sterile saline containing 0.5% heparin.

☒ Sodium Chloride Injection, USP, Preservative-Free, 0.9%, Baxter **Henry Schein Animal Health Catalog #059382**

☒ Heparin Sodium Injection, USP, 10,000 unit/mL **Henry Schein Animal Health Catalog #067792**

A midline abdominal incision was performed to gain access the peritoneum. Three colonic regions of interest - proximal/ascending, transverse, distal/descending - were identified and externalized.

Still-manometry probes (Mikro-CathTM diagnostic pressure catheter, #825-0101, Millar, Houston, TX) were inserted into the colon via a small incision and maintained in position with a loop-hole silk ligature.

Equipment

Mikro-Cath™	NAME
Pressure catheter	TYPE
Millar	BRAND
825-0101	SKU
https://millar.com/clinical/products/mikro-cath#quicktabs-mikro_cath_diagnostic_pressure_c=2	LINK

For the proximal colon, 4 manometry probes were inserted about 10 cm below the ceco-colic junction, at 10, 13, 16 and 19 cm from the point of entry. For the transverse colon, 4 manometry probes were inserted at the end of the proximal colon, at 10, 13, 16 and 19 cm from the point of entry. Distal probes were inserted in the distal colon through the anus with sensors at 10, 13, 16 and 19 cm proximal to the anal verge. One sensor was also added in the anal canal (2 cm from the anal verge).

The abdomen was then closed and the pig placed on his abdomen, leaving access to the back.

The S2-S4 sacral spinal nerves (S1/2 vs S3/4) or T10-L2 thoracolumbar spinal nerves were accessed via a laminectomy. (surgery time: 30 min-40 min).

Laminectomy is a common spine operation and refers to surgical removal of the vertebral lamina, thereby unroofing the vertebral canal. We followed similar processes as described in this [reference](#).

Before surgery, we palpated the lumbar spinous processes along the midline and identified the vertebral levels of interest (T10-L2 or S2-S4). A midline incision using a scalpel/electrocautery was then performed and the subcutaneous tissue and fat dissected to gain access to the thoracolumbar or lumbosacral fascia, respectively. The fascia along the midline was cut to expose the supraspinous ligaments spanning between spinous processes. The supraspinous ligaments was open over a few millimeters. Then using a Freer elevator, the supraspinous ligaments was gently detached from spinous processes (subperiosteal dissection). The dissection was then extended to the interspinous ligaments up to the facet joints of the spinous processes along the area of interest. After locating the lamina, we opened the lamina using a surgical saw or a Kerrison rongeur to extract bone in a piece-wise fashion over the whole vertebral segment of interest in order to gain access to the ligamentum flavum, periosteum, epidural fat, and dura sac. The spinous processes corresponding to the region of interest were removed, allowing access to the meninges. The epidural fat was gently removed taking care not to damage the dura sac and allowing identification of the spinal roots.

During the surgical procedure and as the laminectomy was completed, hemostasis was performed with bipolar electrosurgery. Monopolar electrosurgery was not used because of the proximity of neural structures.

Bone wax was placed along sites of bleeding from exposed bone and absorbable hemostat dressing were used to obtain hemostasis near soft tissue. We also used cotton to wick serous fluid and blood away from the dissection where necessary.

⊗ Absorbable Hemostat Dressing Surgicel® Oxidized Regenerated Cellulose 1/2 X 2 Inch **McKesson Corporation Catalog #188567**

⊗ Bone Wax 2.5 Gram **McKesson Corporation Catalog #2045**

The pigs were euthanized at the end of the experiment with an intravenous injection of pentobarbital (100 mg/kg).

⊗ Euthasol Solution C3N **Henry Schein Animal Health Catalog #009444**

Stimulation experiment

- 3 After surgery, sacral root nerves (S1, S2, S3 and S4) were identified and isolated using small pieces of cotton. Needle electrodes were used to produce stimulation and cuff electrodes were used to apply nerve block either afferent (to assess the role of efferent fibers) or efferent (to investigate the role of afferent fibers).

After positioning the electrodes, recording of the colonic motility using manometry began.

Manometry measurements were made by connecting the still-manometry probes (Mikro-Cath™ diagnostic pressure catheter, #825-0101, Millar, Houston, TX)

Equipment	
Mikro-Cath™	NAME
Pressure catheter	TYPE
Millar	BRAND
825-0101	SKU
https://millar.com/clinical/products/mikro-cath#quicktabs-mikro_cath_diagnostic_pressure_c=2	LINK

via pressure cables (PEC-10D, #850-5090, Millar, Houston, TX)

Equipment	
PEC-10D	NAME
Pressure cable	TYPE
Millar	BRAND
850-5090	SKU
https://millar.com/products/research/pressure/control-units/pcu-2000-dual-channel-pressure-control-unit-patient	LINK

to a transducer (PCU-2000, Millar, Houston, TX)

Equipment	
PCU-2000	NAME
Dual Channel Pressure Control Unit	TYPE
Millar	BRAND
880-0129	SKU
https://millar.com/products/research/pressure/control-units/pcu-2000-dual-channel-pressure-control-unit-patient	LINK

itself connected to a data acquisition system (Micro 1401, CED, Cambridge, UK)

Equipment	
Micro 1401	NAME
Data acquisition unit	TYPE
Cambridge Electronic Design, Ltd	BRAND
CED 1401	SKU
http://ced.co.uk/products/mic3in	LINK

and raw data were collected using the Spike 2 software (CED, Cambridge, UK).

Software

Spike 2

NAME

Windows 7

OS

Cambridge Electronic Design, Ltd

DEVELOPER

A baseline was established for a period of 30 min, then the stimulation was initiated.

The stimulation parameters were as follows: 10 Hz continuous protocol or 30 Hz pulse-train protocol, 0.5 mA, 0.3 ms, 30s-on-90s-off each cycle, 5 cycles.

Following stimulation, recording continued for another 30 minutes.

- 4 Recording of motility data began after surgery was complete and continued for at least 30 min for stabilization of baseline motility, and then for at least another 30 min following completion of the stimulation experiment.

The data in the accompanying dataset was obtained by analyzing the following two subsets of that entire recording: (1) 30 min prior to stimulation; and (2) the 40 minutes immediately following the onset of stimulation (10 min during stimulation, together with another 30 min of recording post stimulation).

The raw manometry data were filtered and rectified using a lab-written Spike 2 code. The same code provided the motility index (MI) defined as the the area under the curve before, during and after stimulation per each minute.