Evaluating the utility of a CO₂ surgical laser for piglet castration to reduce pain and improve wound healing: a pilot study

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1The authors would like to thank personnel at Midwest Veterinary Services for data collection and technical assistance. This project was supported by Agriculture and Food Research Initiative Competitive Grant no. #2020-67015-31540 from the USDA National Institute of Food and Agriculture. Drs. Coetzee and Kleinhenz are supported by the Agriculture and Food Research Initiative Competitive Grants no. #2017-67015-27124, 2020-67030-31479, 2020-67015-31540 and 2020-67015-31546 from the USDA National Institute of Food and Agriculture.

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ABSTRACT

CO₂ surgical lasers are widely used for procedures in veterinary and human medicine. There is evidence to suggest surgery using a CO₂ laser reduces pain and swelling and improves healing time compared to surgery with a scalpel. Millions of piglets in North America are surgically castrated each year using a scalpel. Therefore, piglet welfare may be improved by making refinements to the surgical procedure. The objectives of this preliminary study were to determine the ability of a CO₂ surgical laser to 1) reduce pain and 2) improve wound healing of piglets undergoing surgical castration. Two day old male Yorkshire x Landrace piglets were used and randomly assigned to one of three treatments ($n = 10$ piglets/treatment group): surgical castration with the CO₂ laser, surgical castration with a scalpel, or sham (uncastrated control). Piglets were video recorded in their pens for 1 h pre-procedure and from 0-2, 6-8 and at 24 h post-procedure for behavior scoring. Surgical site images were collected at baseline, 0, 8, 24, 48, 72, 96, 120, 144 and 168 h post-castration for wound healing assessment. Infrared thermography (IRT) images of the surgical site were also taken at baseline, 0, 0.5, 8 and 24 h post-procedure to assess inflammation. Finally, blood was collected from each piglet at baseline and 0.5 h post-castration to assess cortisol levels, prostaglandin E metabolite (PGEM) and pig-major acute phase protein (pig-MAP) concentration. Laser-castrated piglets displayed more pain behaviors across the observation period than scalpel-castrated piglets ($P = 0.05$). Laser-castrated piglets also displayed significantly more agonistic behavior than both scalpel-castrated piglets ($P = 0.005$) and sham piglets ($P = 0.036$); yet, laser-castrated piglets had significantly lower temperatures at the site of incision compared to scalpel-castrated piglets ($P = 0.0211$). There was no
significant difference in wound healing or any of the blood parameters assessed between laser-castrated and scalpel-castrated piglets. There was evidence of thermal tissue damage on the scrotum of piglets that were castrated using the CO$_2$ laser. This may have resulted in the unremarkable healing time and the increased pain behavior observed in this study. The surgical laser technique should be refined before conclusions can be made regarding the utility of a CO$_2$ laser for piglet castration.

**Key words:** animal welfare, castration, CO$_2$ surgical laser, piglet, pain, refinement
ABBREVIATIONS

BW, body weight
SE, standard error
ID, identification
IM, intramuscular
IRT, infrared thermography
MVS, Midwest Veterinary Services
PGEM, prostaglandin E metabolite
Pig-MAP, pig-major acute phase protein
QC, quality control
SNS, sympathetic nervous system
INTRODUCTION

Male piglets in North America are routinely castrated on-farm, to prevent boar taint and minimize aggression (Sutherland, 2015). This painful procedure is done on conscious piglets, by using a scalpel to make an incision on the scrotum and removing the testicles by cutting or tearing the spermatic cord (Rault et al., 2011). Complications, such as hemorrhage, infection, excessive swelling and intestinal herniation (resulting in pre-weaning mortality), may be partially attributable to the described surgical castration technique (Taylor and Weary, 2000; Morales et al., 2017). Refining the castration procedure by replacing the scalpel with a technique that decreases tissue damage and bleeding may reduce these post-surgical complications and lead to improved piglet welfare in commercial production systems.

CO₂ surgical lasers are increasingly being used for procedures in veterinary and human medicine. They function by emitting a colorless, infrared light at a specific wavelength (10,600 nm) which is absorbed by intracellular water and causes tissue cells to ablate or vaporize (Mison et al., 2003). This allows for clean, precise incisions to be made on the skin and takes no more time than if a standard scalpel was used. Pain and swelling have been shown to be significantly reduced in human patients who have undergone the same surgical procedure using a CO₂ laser compared to a scalpel (Tuncer et al., 2010; Lopez-Jornet and Camacho-Alonso, 2013). The CO₂ laser has also refined the canine castration procedure by nearly eliminating blood flow during the surgery and reducing the risk of scrotal hematoma, bruising and infection compared to canine castration with a scalpel (Schultz, 2013).

The objectives of this pilot study were to determine the ability of a CO₂ surgical laser to 1) reduce pain and 2) improve wound healing of piglets undergoing surgical castration. We hypothesized that surgical castration of piglets using the CO₂ laser would result in decreased
inflammation at the surgical site, reduced wound healing time and less post-surgical pain compared to piglets castrated with a scalpel.

**MATERIALS AND METHODS**

All animal use and procedures were approved by the Institutional Animal Care and Use Committee at Midwest Veterinary Services (MVS) prior to study commencement (Protocol # MCL-19065).

**Animals**

Three Yorkshire x Landrace sows nursing one day-old piglets were sourced from MVS Klitz Farm (Oakland, NE) for this study. Sows were examined by a veterinarian and selected based on four parameters: health, body condition, past weaning history and non-aggressive behavior. For enrollment in this study, sows had to be in good physical health, free of any complicating disease, with a body condition score of 3 ± 0.5 out of 5. They also had to have previously weaned at least one litter of piglets (i.e., gilts were excluded). Healthy male piglets with two testicles, intact tails (i.e., no tail docking prior to enrollment), and no palpable hernias were selected from these three sow litters, while the females and males not meeting the selection criteria were cross-fostered to other sows in the unit. Healthy, one day-old male piglets from other litters were then selected and cross-fostered to the study sows until each had 11 piglets (n = 33 piglets total). A record of the biological sows for each of the piglets selected was recorded and is presented in Table 1. Each piglet received two ear tags (one in each ear) with an ID number and were given 1.0 mL of iron dextran (Ferrodex 100 mg/mL; Agri Laboratories, Ltd., St. Joseph, MO) intramuscular (IM) in the neck.

The sows and piglets were transported using a livestock trailer to the Central States Research Center (CSRC; MVS facility, Oakland, NE) 24 h prior to study commencement. Sows and their litters were housed in farrowing crates on raised Tenderfoot® flooring.
Sows had *ad libitum* access to feed and water and were fed a diet that met or exceeded National Research Council (NRC, 2012) nutrient requirements for lactating sows. The facility room temperature was maintained at 24.6 ± 2.5°C, and a heat lamp was provided to each litter of pigs. All animals were exposed to approximately 12 h of light per day.

**Treatments and Processing Procedures**

On the day of the study, piglet weights were collected (mean BW = 1.8 ± 0.7 kg; 2 days old) and their ID numbers were written on their forehead and back using a black permanent marker. This was to aid in piglet identification throughout the study.

Eleven piglets were assigned to each treatment group and treatments were balanced within a litter. Piglets were randomly assigned to one of three treatments: surgical castration using a CO\textsubscript{2} laser (VetScalpel; Aesculight, LLC, Bothell, WA), surgical castration using a scalpel or sham (uncastrated control). Treatment assignments were predetermined by randomizing piglets to treatment group in a spreadsheet (Excel; Microsoft, Redmond, WA) based on piglet and sow ID. One piglet per treatment group (*n* = 3 total) was deemed “extra”, to ensure appropriate study power if a piglet(s) required euthanasia due to post-surgical complications such as inguinal herniation.

To conduct the castration procedure, piglets were removed from their pen, placed on a table in the supine position and restrained by two individuals. The surgical site was then disinfected using gauze soaked in isopropyl alcohol 70% (Vet One; MWI, Boise, ID). Piglets were castrated by making one horizontal incision on the scrotum with the CO\textsubscript{2} laser (set to 15 W, continuous) or scalpel, based on their treatment group. Testicles were removed by ablating (CO\textsubscript{2} laser) or cutting (scalpel) the spermatic cord. The CO\textsubscript{2} laser was calibrated after each litter of pigs to ensure proper functionality (all calibrations yielded 79.7 ± 1.4%; therefore, the actual power of the CO\textsubscript{2} laser at the level of the piglet was 12 W. This is a...
normal deviation for the surgical laser unit). Piglets in the sham treatment group were restrained in the same manner, the handle of the scalpel was used to simulate the incision and the scrotum was manipulated to resemble a surgical castration. Piglets were then returned to their pen. All procedures occurred between 09:30 and 10:30 and were conducted by the same individual with extensive experience in surgically castrating piglets. The length of time piglets were restrained was similar across all treatment groups (approximately 20 sec). Two piglets (one CO₂ laser-castrated and one scalpel-castrated) herniated post-castration and were euthanized using a pentobarbital sodium injection (Fatal-Plus 390 mg/mL; Vortech Pharmaceuticals, Ltd., Dearborn, MI); all other castrated piglets (n = 20) recovered without incident.

**Behavior Recording and Scoring**

Piglets were video recorded for 1 h pre-procedure using a high definition video camera (Sony Handycam HDR-CX405, Sony USA Inc., New York, NY) mounted on a tripod and placed outside of each farrowing pen. After processing, piglets were video recorded for two 3 h periods: 0-2 h post- and 6-8 h post-castration. Finally, 24 h post-procedure, piglets were recorded for 1 h (i.e., 8 h of video data were collected in total for each litter of pigs). The videos were randomized across time point and pen ID using a random number generator (random.org). The behavior of each piglet was scored continuously by one experienced observer for the first 15 mins of every hour of data collected using BORIS software (Behavioral Observation Research Interactive Software v 7.7.3, Torino, Italy) and a detailed ethogram (Table 2). The observer was masked to treatment and time point; however, they could observe which piglets had been castrated and which had not. A total of 3,600 min (60 h) of behavior recordings were scored and analyzed for this study.

Piglet behaviors were analyzed individually and then grouped into categories to assess the activity level of piglets across the observation period and the total proportion of pain...
behaviors displayed. Pain behaviors included tail wagging, trembling, scratching and stiffness (Hay et al., 2003). The active behavior category included walking, running, playing, suckling, nosing and chewing (Viscardi et al., 2018). Inactive behaviors included sleeping and awake inactive.

**Wound Healing**

Still-images of each piglet’s scrotum were collected using a point-and-shoot camera (Olympus Stylus Tough TG-4; Olympus Corporation, Tokyo, Japan), with one individual briefly handling the piglets to facilitate capturing a clear picture of the surgical site. Pictures were taken pre-procedure (baseline) and at 0 h, 8 h, 24 h, 48 h, 72 h, 96 h, 120 h, 144 h and 168 h post-castration. Images were randomized using a random number generator (random.org) and scored using a 6-point scale developed by Sutherland et al., (2010a), by one individual blinded to piglet treatment and time point. Wounds with a score of one were fully healed (no scab) and wounds with a score of six had signs of fresh blood.

**Infrared Thermography (IRT) Imaging**

Infrared thermography images of the surgical castration site were collected from each piglet pre-procedure (baseline) and at 0 h, 0.5 h, 8 h and 24 h post-procedure using a research-grade infrared camera (FLUKE TiX580; FLUKE Corporation, Everett, WA). The camera was calibrated to the ambient temperature and relative humidity of the room prior to taking images. One individual briefly handled the piglets to facilitate image capture while another held the IRT camera in-line with the surgical site at a distance of approximately 0.5 m. In castrated piglets, the incision and surrounding tissues of the scrotum were captured in a single image; in sham piglets, an image of the scrotum was collected. The majority (80%) of IRT data collection coincided with still-image capture of the castration site for wound healing assessment, so piglets were only handled once to minimize stress.
Infrared images were analyzed using research-grade software (SmartView 4.3; FLUKE Corporation, Everett, WA). For each image collected, the temperature of the incision (in castrated piglets) and the average temperature of the surrounding tissues of the scrotum were recorded and analyzed. The difference between the temperatures taken at the two sites on the scrotum was also calculated, by subtracting the temperature of the surrounding tissues by the temperature of the incision. This data was used to assess the degree of inflammation associated with the surgical castration procedure.

**Blood Sample Collection and Processing**

A blood sample (4.0 mL) from each piglet was collected from the jugular vein using a 20-gauge needle (TycoHealth Care, Mansfield, MA) at baseline and 30 min post-castration. Piglets were briefly removed from their pen and restrained in the supine position during sample collection. Blood was immediately transferred into serum separator tubes (BD Vacutainer, Franklin Lakes, NJ) and placed on ice. Once all of the samples at each time point were collected, blood was centrifuged at 3,000 g for 10 min. The serum was pipetted from the tube, placed into cryovials and stored at -80°C until analysis.

Serum samples were submitted to the Iowa State University-Pharmacology Analytical Support Team (ISU-PhAST) at the Iowa State University Veterinary Diagnostic Laboratory for cortisol determination. Samples were also analyzed by a laboratory technician at Kansas State University to determine prostaglandin E metabolite and pig-major acute phase protein concentration. All laboratory personnel were blinded to piglet treatment and time point.

**Cortisol Determination**

Baseline (pre-procedure) and 0.5 h post-procedure serum samples were analyzed for cortisol using the commercially available Cortisol Coated Tube RIA kit (MP Biomedicals catalog no. 07-221105R; Irvine, CA). Samples were run in duplicate. There was a quality control (QC) high and a QC low concentration run to assess drift from beginning to end of
the gamma counter analysis and across runs. The average concentration of the QC low was 48.8 ng/mL and the QC high was 263.7 ng/mL. The intra-assay and inter-assay coefficient of variation was 10.3% and 10.6%, respectively.

**Prostaglandin E metabolite (PGEM) Determination**

Prostaglandin E metabolite (PGEM) was determined from serum samples collected at baseline (pre-procedure) and 0.5 h post-castration. A commercially available PGEM ELISA kit was used (Cayman Chemical catalog no. 514531; Ann Arbor, MI) with minor modifications. Briefly, samples were purified by adding 1.5 mL ice cold acetone to 375 µL serum. Samples were then incubated at -20°C for 30 min followed by centrifugation at 3,000 x g for 5 min. The supernatant was transferred to 13 x 100 mm glass tubes and evaporated using a CentriVap Concentrator (Labconco catalog no. 7810014; Kansas City, MO) and reconstituted with 375 µL of appropriate kit buffer. A 300 µL aliquot of the reconstituted sample was derivatized with proportionally adjusted kit components. Manufacturer protocol was then followed. Samples were diluted 1:5 and ran in duplicate. Absorbance was measured at 405 nm after 60 min of development (SpectraMax i3; Molecular Devices; San Jose, CA). The average concentration of the sample used to determine repeatability across plates was 27.8 pg/mL. The intra-assay and inter-assay coefficient of variation was 16.4% and 19.3%, respectively.

**Pig-Major Acute Phase Protein (pig-MAP) Determination**

Serum samples collected at pre-procedure (baseline) and 0.5 h post-castration were analyzed for pig-MAP using a commercially available pig-MAP ELISA kit (Acuvet Biotech catalog no. AC/PME01; Zaragoza, Spain). Each sample was diluted 1:1,000 and ran in duplicate. Absorbance was read at 450 nm (SpectraMax i3; Molecular Devices; San Jose, CA). The average concentrations used to determine repeatability across plates was 1.6 and
0.8 ug/mL. The intra-assay and inter-assay coefficient of variation was 7.5% and 28.5%, respectively.

**Statistical Analysis**

The total duration of behaviors was converted into proportion of time a piglet engaged in each behavior prior to analysis. This was to account for periods of time when a piglet was not in view and could not be scored. Data were analyzed using a generalized linear mixed (GLIMMIX) model with a beta distribution, including treatment, time, litter, and time x treatment interaction in SAS (Statistical Analysis System 9.4, SAS Institute Inc., NC). Litter was included as a random effect and time was a repeated measure with piglet as the experimental unit. *Post hoc* tests were conducted using the Tukey-Kramer adjustment. Statistical significance was set at *P* < 0.05.

Cortisol was log-transformed for normality prior to analysis. Wound scores, temperature of the surgical site (from IRT images), cortisol, PGEM and pig-MAP were analyzed using a mixed model in SAS, including litter, time, treatment, and time x treatment interaction. Litter was included as a random effect and time was a repeated measure with piglet as the experimental unit. A *post hoc* Tukey’s test was conducted for significant outcomes.

**RESULTS**

**Behavioral Observations**

Piglets were in-view and able to be scored for 90.7 ± 0.05% of the observation period in this study. Four individual behaviors (agonistic: *P* = 0.004, desynchronized: *P* = 0.045, tail wagging: *P* = 0.026, and trembling: *P* = 0.049) and one grouped behavior (pain: *P* = 0.026) were affected by treatment across the observation period (Table 3). Laser-castrated piglets trembled significantly more than scalpel-castrated piglets (*P* = 0.041) and engaged in more desynchronized behaviors (*P* = 0.039). Laser-castrated piglets also wagged their tails
significantly more than sham piglets ($P = 0.027$). Agonistic behavior was displayed significantly more by laser-castrated piglets than both scalpel-castrated ($P = 0.005$) and sham ($P = 0.037$) piglets. Laser-castrated piglets demonstrated significantly more pain behaviors than piglets that were scalpel-castrated ($P = 0.049$).

Eleven individual behaviors and both grouped behaviors (active and pain) were significantly affected by time across the observation period (Table 4): lying ($P = 0.011$), nosing ($P = 0.001$), nosing udder ($P < 0.0001$), sleeping ($P < 0.0001$), standing ($P < 0.0001$), suckling ($P < 0.0001$), walking ($P < 0.0001$), sitting ($P < 0.0001$), tail wagging ($P = 0.003$), desynchronized ($P = 0.009$), agonistic ($P = 0.003$), active ($P < 0.0001$), and pain ($P = 0.032$). Irrespective of treatment group, at 1 h, 2 h, 6 h, and 7 h post-castration, piglets were significantly less active, spending more time lying and sleeping and less time standing and engaged in active behaviors compared to piglets at 0 h, 8 h and 24 h post-procedure ($P < 0.05$). Piglets also spent significantly more time nosing the sow’s udder at 8 h post-castration compared to all other time points ($P < 0.001$). They spent significantly more time nosing other substrates at 24 h post-castration than at 0 h and 8 h ($P < 0.05$). Piglets also spent significantly more time suckling pre-procedure compared to all post-castration time points ($P < 0.0001$). Tail wagging and pain behaviors were not significant after the Tukey-Kramer adjustment. There were no significant behavioral differences between any of the treatment groups pre-castration ($P > 0.05$) and no significant time x treatment effects were found.

**Wound Healing**

There were significant time and treatment effects on wound scores ($P < 0.0001$ for both). Both castrated-treatment groups had significantly higher wound scores than sham piglets ($P < 0.0001$); however, there was no significant difference found in wound score between laser-castrated and scalpel-castrated piglets (Fig.1, $P = 0.988$). Wound scores significantly decreased over time (Fig. 2). In castrated piglets, similar wound scores were
noted during the following post-surgical time ranges: 0 h - 7 h, 7 h - 72 h, 24 h - 120 h, 120 h - 144 h and 144 h - 168 h ($P > 0.05$ for all time ranges). Castration wound scores did not reach baseline levels until 168 h (7 days) post-castration.

A researcher involved in this study, who did not score castration wounds, noted evidence of scrotal tissue burns and bruising of the surrounding tissues in a number of the still-images. Once unblinded to treatment, the images were assessed and both tissue burns and bruising were only observed in piglets that had been castrated using the CO$_2$ laser (Fig. 3).

**Infrared Thermography Imaging**

There were significant time ($P < 0.0001$), treatment ($P = 0.010$) and time x treatment interactions ($P < 0.0001$) found for temperature at the incision site. 0 h post-castration, laser-castrated and scalpel-castrated piglets had significantly higher incision site temperatures compared to sham piglets (Fig. 4a; $P < 0.0001$). At 0.5 h and 7 h post-castration, laser-castrated piglets had significantly lower temperatures at the site of incision compared to sham piglets ($P = 0.0005$ and $P = 0.010$, respectively). Across the assessment period, laser-castrated piglets had significantly lower incision site temperatures compared to scalpel-castrated piglets ($P = 0.008$). Irrespective of treatment, piglets had significantly lower incisional temperatures at 0.5 h post-castration compared to all other time points ($P < 0.0001$).

There was no significant difference in temperature of the surrounding tissues on the scrotum between treatment groups ($P = 0.081$); however, there was a significant effect of time ($P <0.0001$), with piglets having lower temperatures at 0 h and 0.5 h post-castration compared to baseline, 7 h and 24 h.

There were significant time ($P < 0.0001$) and time x treatment interactions ($P < 0.0001$) found for the temperature difference between the two focal sites on the scrotum. The
difference in temperature between the two sites was significant at 0 h post-castration, where both laser-castrated and scalpel-castrated piglets had a 2.8 ± 0.1°C increase in temperature at the site of incision compared to the surrounding tissues (Fig. 4b). At 7 h post-castration, scalpel- and laser-castrated piglets had a 2.0 ± 0.3°C increase in temperature of the surrounding tissues compared to the site of incision. Temperatures of the two sites did not differ significantly between the “incision” and the surrounding tissues on the scrotum of sham piglets.

**Cortisol Concentration**

There were no treatment ($P = 0.19$) or treatment x time interactions ($P = 0.59$) found for plasma cortisol concentrations. There was a significant time effect observed ($P = 0.01$). Cortisol concentrations were elevated for both surgical castration groups at 0.5 h. Mean plasma cortisol concentrations pre-castration were 447.0 ± 55.7 ng/mL, 389.9 ± 63.2 ng/mL and 582.1 ± 137.1 ng/mL for the laser-castrated, scalpel-castrated and control piglets, respectively (Table 5). At 0.5 h after castration, the cortisol concentrations were 659.2 ± 69.0 ng/mL, 540.7 ± 67.5 ng/mL and 588.9 ± 83.1 ng/mL for the laser-castrated, scalpel-castrated and control piglets, respectively.

**Prostaglandin E metabolite Concentration**

There were no treatment ($P = 0.62$), time ($P = 0.61$) or treatment x time interactions ($P = 0.84$) for PGEM concentrations. The PGEM concentrations prior to castration were 157.1 ± 24.9 pg/mL, 153.1 ± 36.2 pg/mL and 139.7 ± 13.4 pg/mL for the laser-castrated, scalpel-castrated, and control piglets, respectively. At 0.5 h after castration, the PGEM concentrations were 157.8 ± 20.0 pg/mL, 129.8 ± 17.7 pg/mL and 135.5 ± 16.1 pg/mL for the laser-castrated, scalpel-castrated and control piglets, respectively.
**Pig-Major Acute Phase Protein Concentration**

No treatment \((P = 0.73)\), time \((P = 0.62)\) or treatment x time interactions \((P = 0.92)\) were observed for the pig-MAP concentrations. Mean pig-MAP concentrations pre-castration were \(0.97 \pm 0.07 \text{ mg/mL}, 0.94 \pm 0.10 \text{ mg/mL} \) and \(0.88 \pm 0.09 \text{ mg/mL}\) for the laser-castrated, scalpel-castrated and control piglets, respectively. At 0.5 h post-castration, pig-MAP concentrations were \(0.96 \pm 0.10 \text{ mg/mL}, 0.91 \pm 0.11 \text{ mg/mL} \) and \(0.87 \pm 0.06 \text{ mg/mL}\) for the laser-castrated, scalpel-castrated and control piglets, respectively.

**DISCUSSION**

This study examined the ability of a CO\(_2\) surgical laser to reduce pain and improve wound healing of piglets undergoing surgical castration. Piglets that were castrated using the CO\(_2\) laser exhibited significantly more pain behaviors (trembling, spasms, rubbing the rump, tail wagging and stiffness) than scalpel-castrated piglets, which is contrary to the study hypothesis. Three factors likely contributed to this result: the disinfectant used, piglet restraint and laser-castration technique. Isopropyl alcohol 70% is a common disinfectant used in human and veterinary medicine. Alcohols are highly flammable in nature (CDC, 2008) and there is evidence that heat produced by the CO\(_2\) laser caused a chemical reaction with the alcohol on the piglet’s scrotal surface, resulting in thermal tissue damage. In humans, moderate or severe pain after a thermal burn is common (Perry et al., 1981; McIntyre et al., 2016) and this may explain the increased pain behavior observed in laser-castrated piglets. Based on case reports in the human medicine literature, pooling of alcohol-based skin preparations should be avoided, and the disinfectant used should be dried completely (i.e., allowed to evaporate) when using a surgical laser, to prevent burns (Tooher et al., 2004; Jones et al., 2017). In this study, castration of piglets occurred immediately after disinfecting the surgical site, to limit the stress associated with piglet restraint and separation from the sow. Rather than use an alcohol-based disinfectant and wait the recommended drying time (2-
3 mins) with piglets restrained, a nonalcohol-based solution could be used in future work with a CO$_2$ laser (Vo and Bengezi, 2014; Jones et al., 2017). Difficulty in completely immobilizing conscious piglets resulted in suboptimal laser-castration technique, which also likely contributed to the increase in pain behavior observed. Improving piglet restraint to restrict movement during the castration procedure will allow for a more precise incision to be made with the CO$_2$ laser and prevent poor technique from confounding pain behavior results. Laser-castrated piglets engaged in more agonistic behavior across the observation period than all other study piglets. Pain increases aggressive behavior, irritability and negatively affects interpersonal relations in animal species, including dogs, cats, rats and horses (Curtis, 2008; Fureix et al., 2010; Camps et al., 2012; Njoku et al., 2015); therefore, it could be anticipated that an increase in pain behavior correlated with an increase in agonistic behavior in this study.

There was no difference in wound score (i.e., healing time) between scalpel-castrated and laser-castrated piglets throughout the study. This is likely related to the burning of the scrotal tissue in laser-castrated piglets. When a surgical laser causes thermal tissue damage, a slower healing time within the first few days or weeks has been observed (dogs: Durante and Kriek, 1993; Mison et al., 2003; pigs: Buell and Schuller, 1983; Molgat et al., 1994; Schoinohoriti et al., 2012). The lack of difference in healing time when comparing a CO$_2$ laser to a scalpel may also be a result of poor laser technique or a power setting that was too low, requiring multiple passes with the laser to make an incision (Reid, 1991; Capon and Mordon, 2003). These sequential passes with the laser at the incisional site increases the amount of thermal injury (Reid, 1991). The power setting selected for this study (15 W, continuous) was determined from the manufacturer’s recommendations and by practicing surgical castration on piglet cadavers prior to study start. The power may need to be increased in future work, as multiple passes with the laser were required to make the scrotal incision.
Infrared thermography is a validated tool to measure cutaneous temperature and assess inflammation (Calkosinski et al., 2015; Soroko and Howell, 2018). Laser-castrated piglets had a lower temperature at the incision site across the assessment period (up to 24 h post-castration) compared to scalpel-castrated piglets, suggesting that there was less inflammation at the surgical site when the CO₂ laser was used. While this result is consistent with our study hypothesis, it contradicts the literature regarding thermal tissue injury and inflammation (Strudwick and Cowin, 2017). As well, a decrease in inflammation after piglet surgical castration is generally associated with a reduction in acute pain and pain behaviors (Herskin and Di Giminiani, 2018), yet this was not observed. The temperature of the surrounding scrotal tissues (excluding the incision site) did not differ between laser-castrated and scalp-castrated piglets, suggesting inflammation at this location may be more predictive of castration-associated pain. The significant drop in temperature from 0 h to 0.5 h post-castration may be related to activation of the sympathetic nervous system (SNS) in piglets, leading to peripheral vasoconstriction and a decrease in cutaneous temperature. This has been observed in animals that are stressed or in pain (Stewart et al., 2005; Dockweiler et al., 2013; Bates et al., 2014). At 0.5 h post-castration, piglets were separated from the sow, restrained, and a blood sample was collected via jugular venipuncture, all prior to collecting an IRT image. The amount of handling, restraint and painful procedures piglets in this study were subjected to within a 30 mins period almost certainly caused stress and acute physiological changes. Future work should prioritize noninvasive data collection (e.g., IRT, behavior…etc.) prior to conducting more invasive procedures (e.g., venipuncture) to minimize stress and SNS activation from potentially confounding results.

Blood cortisol has been widely used as a biomarker of stress and pain in animals. In response to surgical castration, cortisol levels increase in piglets and peak 30-90 mins after processing (Prunier et al., 2005; von Borell et al., 2009; Bates et al., 2014; Tenbergen et al.,
The mean plasma cortisol concentration of piglets in this study significantly increased from baseline to 0.5 h post-castration; however, there was no effect of treatment (i.e., sham piglets who were not castrated also had a significant increase in blood cortisol level). While this contradicts previous work where cortisol levels did not significantly increase in piglets that were handled only and not subjected to a painful procedure (Prunier et al., 2005; Carroll et al., 2006; Marchant-Forde et al., 2014), it is important to note that cortisol is not a specific biomarker of pain. Stress from handling or restraint can cause an increase in plasma cortisol concentration (Llamas Moya et al., 2008). Many piglets in this study were cross-fostered and all were transported to the research facility less than 24 h prior to study start. Both of these events are known to be stressful (Sutherland et al., 2010b; Calderón Diaz et al., 2018). Acclimation of research animals to a new environment is ideal, to reduce the risk of stress interfering with the study results.

Prostaglandin E and acute phase proteins in the blood increase in response to stress, pain and inflammation (Piñeiro et al., 2013; Davidson et al., 2014). In this study, there were no treatment or time differences in these outcome measures (PGEM and pig-MAP), which may relate to the high levels of stress piglets were already experiencing due to the recent cross-fostering and transportation events.

Using a CO₂ surgical laser instead of a scalpel has the potential to reduce pain, inflammation and improve animal welfare (Tuncer et al., 2010; Lopez-Jornet and Camacho-Alonso, 2013). In this study, thermal tissue damage caused by the CO₂ laser confounded the pain and wound healing results, making it difficult to draw conclusions regarding the utility of this tool for surgical castration of piglets. A nonalcohol-based disinfectant should be used and the laser-castration technique optimized in future work, to more accurately assess pain, inflammation and wound healing in piglets after processing.
DISCLOSURES

The authors declare that there is no conflict of interest.
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FIGURE LEGENDS

**Fig. 1:** Average wound score (±SE) of piglets in each treatment group over time. Asterisks represent a significant difference ($P < 0.05$) between the castrated piglets (laser and scalpel; $n = 20$) and sham piglets ($n = 10$).

**Fig. 2:** Average wound score (±SE) of castrated piglets ($n = 20$) over time. Different letters indicate significance ($P < 0.05$).

**Fig. 3:** A comparison of the surgical wound at 24 h post-castration for a piglet undergoing the procedure using **a)** the CO$_2$ laser or **b)** a scalpel. Evidence of tissue burning at the incision site and bruising of the surrounding tissues is clear in the laser-castrated piglet.

**Fig. 4:** Average temperature ($^\circ$C) **a)** at the incision site (±SE) and **b)** temperature difference (±SE) between the incision site and the surrounding tissues of the scrotum of piglets in each treatment group over time. Asterisk represent a significant difference ($P < 0.05$) between the castrated piglets ($n = 20$) and sham piglets ($n = 10$); letter represents a significant difference between laser-castrated piglets ($n = 10$) and sham piglets.
**Table 1**: Record of male piglets and their biological sow after cross-fostering.

| Piglet ID | Biological Sow ID | Study Sow ID | Pen # in Research Facility |
|-----------|-------------------|--------------|---------------------------|
| 1         | 17138             | 16176        | 1                         |
| 2         | 18063             |              |                           |
| 3         | 18063             |              |                           |
| 4         | 18063             |              |                           |
| 5         | 16176             |              |                           |
| 6         | 16176             |              |                           |
| 7         | 16176             |              |                           |
| 8         | 16176             |              |                           |
| 9         | 16176             |              |                           |
| 10        | 16176             |              |                           |
| 11        | 16176             |              |                           |
| 12        | 16177             | 18070        | 3                         |
| 13        | 16177             |              |                           |
| 14        | 16177             |              |                           |
| 15        | 16177             |              |                           |
| 16        | 18070             |              |                           |
| 17        | 18070             |              |                           |
| 18        | 18070             |              |                           |
| 19        | 18070             |              |                           |
| 20        | 18070             |              |                           |
| 21        | 18070             |              |                           |
| 22        | 18070             |              |                           |
| 23        | 16114             | 18159        | 2                         |
| 24        | 16114             |              |                           |
| 25        | 16114             |              |                           |
| 26        | 16114             |              |                           |
| 27        | 16114             |              |                           |
| 28        | 16114             |              |                           |
| 29        | 18159             |              |                           |
| 30        | 18159             |              |                           |
| 31        | 18159             |              |                           |
| 32        | 18159             |              |                           |
| 33        | 18159             |              |                           |
Table 2: Ethogram used to score piglet behavior, grouped into feeding, locomotion, non-specific behaviors, castration-related pain behaviors, posture and social cohesion (adapted from Hay et al., 2003).

| Behavior     | Description                                                                 |
|--------------|-----------------------------------------------------------------------------|
| Suckling     | Teat in mouth and suckling movements                                         |
| Nosing udder | Nose in contact with udder, up and down head movements                       |
| Playing      | Springing, bouncy movements with or without littermates                      |
| Agonistic    | Biting or fighting other littermates                                          |
| Walking      | Moving forward at a normal pace                                              |
| Running      | Trot or gallop                                                              |
| Awake inactive | No special activity, but awake                                             |
| Sleeping     | Lying down, eyes closed                                                     |
| Nosing       | Snout in contact with a substrate                                           |
| Chewing      | Nibbling at littermates or substrates                                        |
| Trembling    | Shivering, as with cold                                                     |
| Spasms       | Quick and involuntary contractions of the muscles                           |
| Scratching   | Rubbing the rump against the floor, pen walls, or littermates               |
| Tail wagging | Tail’s movement from side to side (or up and down)                          |
| Stiffness    | Lying with extended and tensed legs                                         |
| Lying        | Body weight supported by side or belly                                      |
| Sitting      | Body weight supported by hindquarters and front legs                        |
| Standing     | Body weight supported by four legs                                          |
| Kneeling     | Body weight supported by front carpal joints and hind legs                  |
| Isolated     | Alone, or with one littermate, distance of 40 cm separates the animal(s) from the closest group of littermates |
| Desynchronized | Activity different from that of most littermates (at least 75%)              |
Table 3: Proportion of time piglets were engaged in specific behaviors (n = 10 piglets per treatment group) post-castration. Values represent the proportional means (± SE).

| Behavior² | Post-Castration Treatment P-value | CO₂ Laser | Scalpel | Sham |
|-----------|----------------------------------|-----------|---------|------|
| Tail wagging | 0.0257 | 0.02±0.00ᵃ | 0.00±0.00ᵇ | 0.00±0.00ᵇ |
| Trembling | 0.0493 | 0.07±0.07ᵃ | 0.02±0.02ᵇ | 0.04±0.05ᵇ |
| Desynchronized | 0.0446 | 0.18±0.07ᵃ | 0.05±0.03ᵇ | 0.16±0.06ᵇ |
| Agonistic | 0.0038 | 0.01±0.00ᵃ | 0.00±0.00ᵇ | 0.00±0.00ᵇ |
| Pain² | 0.0257 | 0.06±0.01ᵃ | 0.02±0.00ᵇ | 0.03±0.01ᵇ |

¹Only significant behavior variables are presented
²Pain behaviors include: scratching, stiffness, trembling, tail wagging
ᵃᵇValues within a row with different superscripts are significantly different (P < 0.05)
Table 4: Proportion of time piglets were engaged in specific behaviors (n = 30 piglets total) pre- and post-castration. Values represent the proportional means (± SE).

| Behavior  | Pre-Castration | Post-Castration | 0h | 1h | 2h | 6h | 7h | 8h | 24h |
|-----------|----------------|----------------|-----|----|----|----|----|----|-----|
| Baseline  | 0.71±0.03      | 0.0107         | 0.47±0.03 | 0.84±0.03 | 0.94±0.03 | 0.79±0.03 | 0.93±0.02 | 0.52±0.03 | 0.42±0.04 |
| Time P-value | <.0001         | <.0001         | 0.02±0.01 | 0.03±0.01 | -   | 0.00±0.03 | 0.02±0.02 | 0.02±0.00 | 0.10±0.02 |
| Lying     | 0.07±0.02      | 0.02±0.01      | 0.27±0.05 | 0.15±0.04 | 0.13±0.04 | 0.18±0.04 | 0.11±0.04 | 0.41±0.06 | 0.20±0.04 |
| Nosing    | 0.17±0.04      | 0.02±0.02      | 0.44±0.04 | 0.14±0.03 | 0.02±0.02 | 0.19±0.03 | 0.09±0.02 | 0.12±0.03 | 0.50±0.03 |
| Nosing udder | 0.52±0.11      | <.0001         | 0.03±0.02 | 0.03±0.02 | -   | 0.08±0.03 | -    | 0.11±0.04 | 0.17±0.06 |
| Sleep     | 0.05±0.02      | <.0001         | 0.15±0.02 | 0.06±0.02 | 0.00±0.01 | 0.02±0.00 | 0.05±0.01 | 0.07±0.01 | 0.10±0.02 |
| Standing  | 0.13±0.02      | <.0001         | 0.12±0.02 | 0.04±0.02 | 0.07±0.03 | 0.03±0.01 | 0.06±0.02 | 0.17±0.02 | 0.09±0.02 |
| Suckling  | 0.00±0.00      | 0.0030         | 0.01±0.00 | 0.00±0.00 | 0.01±0.01 | 0.02±0.00 | 0.01±0.00 | 0.00±0.00 | 0.03±0.00 |
| Walking   | 0.12±0.03      | 0.0086         | 0.26±0.07 | 0.04±0.02 | 0.40±0.16 | 0.13±0.09 | 0.01±0.03 | 0.18±0.07 | 0.10±0.02 |
| Tail wagging | 0.29±0.03      | <.0001         | 0.53±0.04 | 0.16±0.03 | 0.06±0.03 | 0.21±0.03 | 0.07±0.02 | 0.48±0.04 | 0.57±0.04 |

Only significant behavior variables are presented

Active behaviors include: nosing, suckling, walking, chewing, playing, running

Pain behaviors include: scratching, stiffness, trembling, tail wagging

Dash indicates behavior was not observed

Not significant after Tukey-Kramer adjustment

Values within a row with different superscripts are significantly different (P < 0.05)
Table 5: Mean (95% confidence interval) analyte concentration of cortisol, pig-MAP\(^1\), and PGEM\(^2\) analyzed for piglets undergoing surgical castration by CO\(_2\) laser, scalpel or sham (\(n = 10\) piglets per treatment group).

|          | CO\(_2\) Laser | Scalpel | Sham   | Treatment | Time | Treatment x time |
|----------|----------------|---------|--------|-----------|------|------------------|
| **Cortisol, ng/mL** |                |         |        |           |      |                  |
| Baseline | 447.0          | 389.9   | 582.1  |           |      |                  |
|          | 321.1 – 573.0  | 246.9 – 532.9 | 271.9 – 892.4 | 0.19 | 0.01 | 0.59             |
| 0.5 h    | 695.2          | 540.7   | 588.9  |           |      |                  |
|          | 534.8 – 855.7  | 380.2 – 701.2 | 428.4 – 749.3 |      |      |                  |
| **Pig-MAP, mg/mL** |                |         |        |           |      |                  |
| Baseline | 0.97           | 0.94    | 0.88   |           |      |                  |
|          | 0.71 – 1.23    | 0.69 – 1.20 | 0.62 – 1.14 | 0.73 | 0.62 | 0.92             |
| 0.5 h    | 0.96           | 0.91    | 0.87   |           |      |                  |
|          | 0.71 – 1.22    | 0.65 – 1.16 | 0.61 – 1.13 |      |      |                  |
| **PGEM, pg/mL** |                |         |        |           |      |                  |
| Baseline | 157.1          | 153.1   | 139.7  |           |      |                  |
|          | 114.6 – 199.7  | 110.5 – 195.7 | 97.0 – 182.3 | 0.62 | 0.61 | 0.84             |
| 0.5 h    | 157.8          | 129.8   | 135.5  |           |      |                  |
|          | 115.2 – 200.3  | 87.2 – 172.4 | 92.9 – 178.1 |      |      |                  |

\(^1\)Pig-MAP: Pig-major acute phase protein  
\(^2\)PGEM: Prostaglandin E metabolite
Figure 2

Average Wound Score

Baseline 0 h 7 h 24 h 48 h 72 h 96 h 120 h 144 h 168 h

Time Post-Castration (Hour)

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Figure 3
Figure 4

(a) Average Temperature, °C

(b) Average Temperature Difference, °C

Legend:
- Laser
- Scalpel
- Sham

Time Post-Castration, Hour

Baseline 0 h 0.5 h 7 h 24 h