

# The effects of handling and group size on welfare of pigs in lairage and their influence on stomach weight, carcass microbial contamination and meat quality

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Rabaste, C., Faucitano, L., Saucier, L., Mormède, P., Correa J. A., Giguère, A. and Bergeron, R. 2007. **The effects of handling and group size on the welfare of pigs in lairage and their influence on stomach weight, carcass microbial contamination and meat quality.** *Can. J. Anim. Sci.* **87**: 3–12. At unloading and on the way to stunning, 800 barrows were exposed to either gentle handling (GH: slowly with a plastic board or whip) or rough handling (RH: quickly with an electric prod). Pigs were kept in large or small groups (30 or 10 pigs) during lairage. Compared with GH, RH increased climbing ( $P < 0.05$ ), slipping ( $P < 0.01$ ) and turning around ( $P < 0.001$ ) behaviours during unloading, and climbing ( $P < 0.05$ ) on the way to stunning. RH also reduced drinking behaviour during lairage ( $P < 0.01$ ). Pigs kept in large groups were observed more often standing ( $P < 0.05$ ) and fighting ( $P < 0.001$ ) than pigs kept in small groups, but, in contrast, had a slightly lower level of urinary cortisol at slaughter. Stomach weight and microbial contamination at slaughter were not affected by treatments. RH tended to increase skin bruise score on the carcass ( $P < 0.06$ ) and produced more exudative meat ( $P < 0.05$ ). In conclusion, the response of pigs to the two specific stressors applied prior to slaughter in this study did not seem to contribute to stomach weight variation at slaughter, but it did influence pork quality.

**Key words:** Pigs, pre-slaughter handling, group size, stress, stomach weight, microbial contamination, behaviour, meat quality

Rabaste, C., Faucitano, L., Saucier, L., Mormède, P., Correa J. A., Giguère, A. et Bergeron, R. 2007. **Effets de la manipulation et de la taille du groupe sur le bien-être des porcs à l'abattoir et leur influence sur le poids des estomacs, la contamination microbienne de la carcasse et la qualité de la viande.** *Can. J. Anim. Sci.* **87**: 3–12. Au déchargement et en route vers l'anesthésie, 800 porcs mâles castrés ont été soumis à des manipulations douces (GH: lentement avec un panneau de plastique ou un fouet) ou rudes (RH: rapidement avec un aiguillon électrique). Pendant la période de repos, les porcs ont été maintenus en grands ou en petits groupes (30 vs. 10). Les RH, comparés aux GH, ont augmenté au déchargement les chevauchements ( $P < 0,05$ ), les glissades ( $P < 0,01$ ) et les demi-tours ( $P < 0,001$ ) et sur la route vers l'anesthésie, les chevauchements ( $P < 0,05$ ). Les RH ont également réduit les abreuvements pendant l'attente ( $P < 0,01$ ). Les porcs maintenus en grands groupes ont été plus souvent observés debout ( $P < 0,05$ ) et en combat ( $P < 0,001$ ) que les porcs maintenus en petits groupes, mais en revanche, ils ont eu une concentration de cortisol urinaire légèrement plus faible à l'abattage. Le poids des estomacs et la contamination microbienne à l'abattage n'ont pas été affectés par les traitements. Les RH ont eu tendance à avoir un score de lésions sur la peau plus élevé ( $P < 0,06$ ) et à produire des viandes plus exsudatives ( $P < 0,05$ ). En conclusion, la réponse des porcs aux deux facteurs de stress spécifiques appliqués avant l'abattage dans cette étude n'a pas semblé contribuer à la variation du poids des estomacs, mais a influencé la qualité de la viande.

**Mots clés:** Porcs, manipulation pré-abattage, taille du groupe, stress, poids des estomacs, contamination microbienne, comportement, qualité de la viande

Slaughtering animals with a full stomach is considered a high risk factor for meat safety, as spillage of gut contents, due to more frequent inadvertent puncture of the stomach during the dressing process, can lead to microbial cross-contamination between carcasses (Miller et al. 1997). To reduce the risk of puncturing the stomach, a feed withdrawal of 16 to 24 h before slaughter has been recommended to reduce

stomach size (Chevillon 1994). However, industry reports and some studies have revealed a high variability in stomach weights at slaughter, even among pigs that were subjected to the same fasting interval before slaughter (Guise et al. 1995; Turgeon 2003). According to Enck et al. (1989), stress

**Abbreviations:** CA, catecholamines; DL, drip loss; DM, dry matter; EC, electrical conductivity; GC, glucocorticoids; LT, Longissimus thoracis; RH, rough handling; GH, gentle handling; TAM, total aerobic mesophilic

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increases intestinal motility, resulting in a greater evacuation of the caecum and large intestine. It also increases the pH of the stomach contents, favouring the survival, proliferation and release of faecal bacteria (such as *Salmonella*), both towards the internal organs and the surrounding environment (Gregory 1998). Hence, the individual difference in the pig response to preslaughter stress might contribute to stomach weight variation at slaughter.

Besides creating a reservoir of animals aimed at maintaining the constant speed of the slaughter line, the function of lairage is to allow the animals to recover from the stress endured during transport and unloading, and consequently to improve meat quality. Hence, a lairage time of 2 to 4 h is recommended (Warriss 2003). However, these benefits can be lost if lairage facilities are not well designed and poor handling procedures are applied (Hunter et al. 1994; Faucitano et al. 1998; Van der Wal et al. 1999). The use of electric prods must be limited in pig handling given their detrimental effects on welfare (flight behaviour, higher heart rate and salivary cortisol level) and meat quality (Brundige et al. 1998; D'Souza et al. 1998; Jongman et al. 2000). However, the high slaughter speed of modern abattoirs is often assured by the indiscriminate use of this handling device (Geverink et al. 1996; Jones 1999).

Mixing unfamiliar pigs inevitably causes some fighting, which causes skin bruises and poor pork quality (Jones et al. 1994; Warriss 1996). However, it is a common practice to mix large groups of pigs (up to 90 per pen) either because of the lack of adequate holding facilities (adjustable pen size) or because a change in this practice is not perceived by the abattoir managers as economically important. To limit fighting and help pigs rest and recover from transport stress, the current recommendations are either to keep pigs in small groups (10 to 15 pigs) or to mix very large groups (up to 200 pigs) in the lairage pen (Grandin 1990; Christensen and Barton-Gade 1997). However, as emphasized by Warriss (2000), the degree of fighting during lairage is rather related to the harshness of previous handling than on group size. Turner et al. (1999) showed that group size influences drinking behaviour, pigs in large groups (60) spending significantly less time drinking than pigs in smaller groups (20). Hence, it may be hypothesized that the difference in drinking rate between groups of pigs varying in size may contribute to stomach weight variation at slaughter.

The aim of this experiment was thus to determine, under commercial conditions, the effects of gentle vs. rough handling practices and large vs. small group size in lairage on behaviour, stomach weight, microbial carcass contamination and meat quality variation of pigs.

## MATERIALS AND METHODS

### Animals and Treatments

Ten batches of 80 crossbred barrows (synthetic line provided by F. Ménard, Ange-Gardien, QC; average weight: 108.1 ± 4.1 kg) were randomly selected, identified with an individual mark on their back and side at the same farm (one batch per week), fasted (16 h prior to loading) and transported in separate compartments of 20 pigs each for 1 h to a

commercial abattoir. At unloading, pigs from each batch were randomly distributed between two handling treatments (RH: rough and GH: gentle), and two group sizes (10 and 30) were formed in the lairage pens. Pigs handled roughly were moved as quickly as possible with an electric prod, whereas pigs handled gently were moved slowly with a plastic board during unloading, and a whip (used to tap on the back, only when necessary) on their way to the restrainer. Pigs handled roughly each received one electric shock upon unloading from the truck, when entering the lairage pen, and on their way to the restrainer (Fig. 1). During unloading and when moved towards the restrainer, all animals were handled in groups of 10.

In lairage, pigs from both large and small groups were kept at a stocking density of 0.59 m<sup>2</sup>/pig per pen, with a ratio of one water nipple per 10 pigs. Pigs were showered for 13 min from the start of the lairage period, and for 15 min before being slaughtered. All pigs were kept in lairage for 3 h and fasted for a total of 20 h. Animals were cared for according to the recommended codes of practice for swine (Agriculture and Agri-Food Canada 1993) and to the guidelines of the Canadian Council on Animal Care (1993), with the exception of the use of a whip in the stunning raceway, which reflects commercial practice.

### Behavioural Recordings

During unloading from the truck, the frequency of climbing (escape behaviour), slipping and turning around was recorded on all groups of 10 pigs belonging to the small group size treatment, and on randomly selected subgroups of 10 pigs from each large group size treatment (total sample size of 400 pigs). During the first hour of lairage, standing, lying down, drinking and aggression were recorded over four periods of 8 min on the same animals. Aggressive behaviours were classified as fighting and agonistic acts (bites and head knocks). The number of pigs standing and lying down was recorded by scan sampling at 2-min intervals, while the frequency of drinking and aggressive behaviours was recorded continuously. When pigs were moved to the restrainer, the frequency of climbing and turning around was recorded, but only on pigs that had been kept in groups of 10 during lairage. Behavioural variables were noted by two observers placed along the resting pens and the raceways. Over the experiment, the observers alternated between treatment combinations for a more reliable behavioural evaluation.

### Physiological Measures

Urine samples were collected directly on the slaughter line from the bladders of the same animals observed during lairage. Following the addition of HCl 6 M (1% of urine volume) as preservative, samples were immediately frozen (-45°C) pending analysis of urinary free corticoids (cortisol and cortisone) and catecholamines (CA) (noradrenalin, adrenaline and dopamine). Corticoids were measured by high pressure liquid chromatography (HPLC) coupled with a UV detector after extraction on reverse phase columns (Hay and Mormède 1997a). CA were measured by HPLC coupled with an electrochemical detector after extraction on

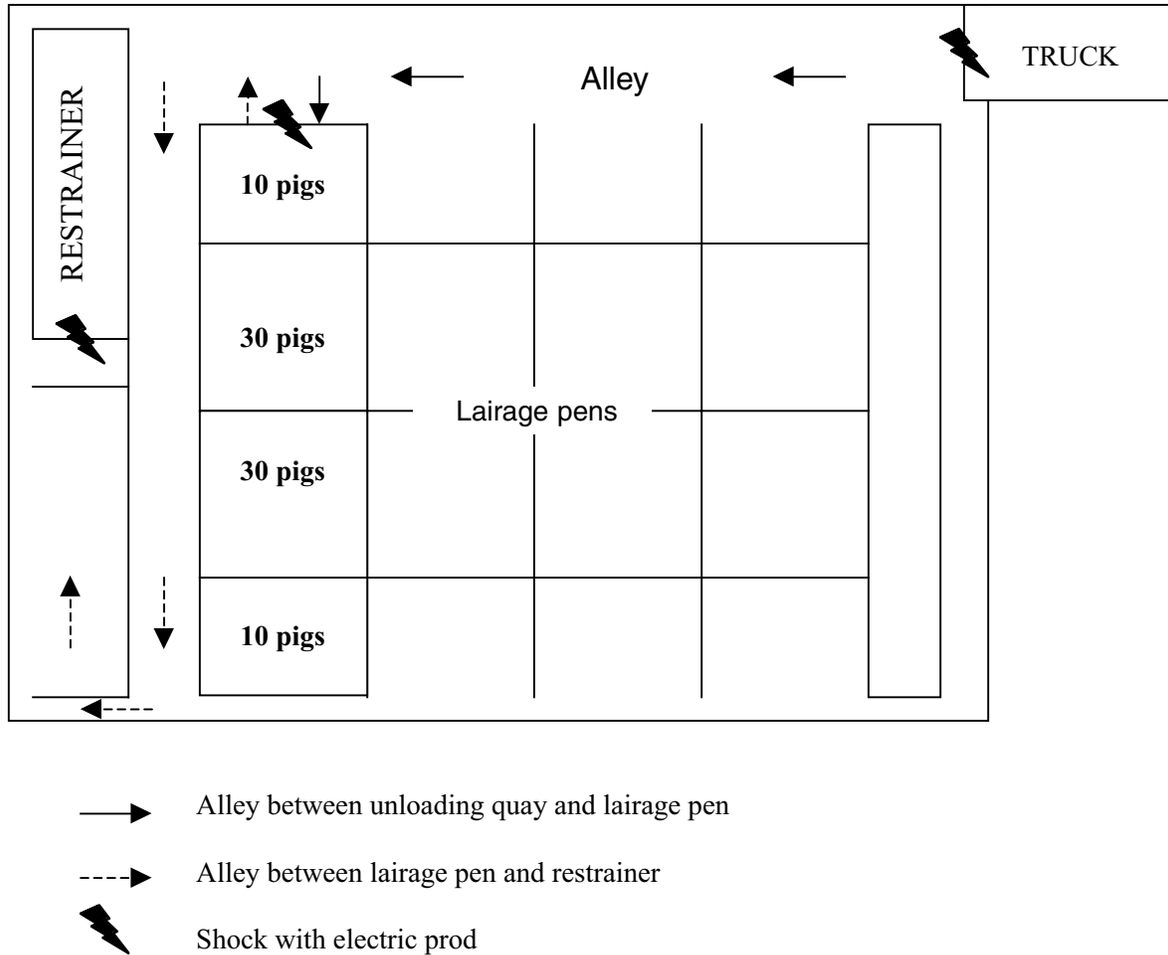


Fig. 1. Plan of the lairage area and distribution of the treatments.

cationic columns (Hay and Mormède 1997b). Creatinine was measured by a colorimetric quantitative method (Procedure 500, Sigma Diagnostics, Saint-Quentin-Fallavier, France) to correct for urine dilution.

### Stomach Weight and Content Composition

Stomachs were removed from 400 carcasses on the dressing line. They were identified and stored at 4°C until they were weighed, full and emptied of their content. Stomach content weights were expressed on a wet weight basis and the state of fasting was assessed according to the criteria set by Chevillon (1994). The contents were mixed and a qualitative assessment of contents was made (sawdust from the truck, liquid or feed). The percentage of dry matter (DM) was calculated after lyophilization at 50°C.

### Microbial Analysis on the Carcass

Carcasses from all animals that had been observed during unloading and lairage were sampled. Swabbing was carried out using a sterile sponge kept in a Whirl-pak™ sampling bag (#B01245, Nasco, Fort Atkinson, WI). Ten milliliters of

diluent constituted of 0.1% peptone water supplemented with 1% Tween 80 was added to each bag and samples were maintained on ice for transportation and kept at 4°C until analysis. The sponge was applied on the internal rib cage, on the briskets and on the top of the two front feet for a total approximate surface of 983 cm<sup>2</sup>. *Escherichia coli*, coliforms and total aerobic mesophilic (TAM) counts were performed using hydrophobic grid membrane techniques described by Gill and Jones (2000) using the Spreadfilter from Filtaflex Ltd. (Almonte, ON). Samples were stomached for 2 min prior to cell enumeration. Serial dilutions were filtered and membranes (ISO-GRID™, Neogen, Lansing, MI) were incubated on appropriate agar medium. TAM counts were performed on Tryptic Soy Agar (TSA; Becton Dickinson, Mississauga, ON) incubated at 35°C for 48 h and colonies were stained with a 0.1% solution of Triphenyltetrazolium chloride (Sigma Aldrich, Oakville, ON) for 15 min. Coliforms were enumerated on Lactose Monesin Glucuronate agar (LMG; QA Life Sciences, San Diego, CA) after an incubation of 24 h at 35°C. The filters were then transferred on buffered 4-methylumbeliferoyl-β-D glu-

curonide agar (BMA; QA Life Sciences) and incubated for 2 h at 35°C for *E. coli* enumeration after illumination of the grid with a long-wave UV light. Grids were photographed (Coolpix995, Nikon Corp., Tokyo, Japan) and the most probable number (MPN) was determined using the software HGMF Interpreter (Filtaflex Ltd). Pieces of meat trimmed off on the processing line for visual contamination were recorded for each carcass.

### Skin Bruises and Meat Quality Measurements

Skin bruises were assessed in the cooler immediately after dressing using a photographic scale from 0 (none) to 5 (severe; MLC 1985). Muscle pH was assessed at 45 min (pH<sub>1</sub>) and 24 h (pHu) *post mortem* with a pH meter (Oakton Instruments Model pH 100 Series, Niles, IL) fitted with a Cole Parmer spear type electrode (Cole Palmer Instrument Company, Vernon Hills, IL) and an automatic temperature compensation probe by insertion in the *longissimus thoracis* (LT) muscle between the 3rd and 4th last ribs (Canadian grading site). At 24 h post mortem the following measures were taken from the LT muscle in the carcass grading site region: electrical conductivity (EC) by insertion of the Pork Quality Meat probe (PQM-I-INTEK, GmbH, Aichach, Germany), subjective colour score using the Japanese Color Standards (JCS) ranging from 1 = pale to 6 = dark color (Nakai et al. 1975), light reflectance using a Minolta Chromameter CR 300 (D65 light source with 0° viewing angle geometry) according to the reflectance coordinates (CIE  $L^*$ ,  $a^*$ ,  $b^*$ ), and after exposing the muscle surface during a 1-h blooming time. Drip loss (DL) was assessed on an adjacent chop according to a modified “juice container” procedure (Correa et al. 2007). Briefly, three cylindrical muscle cores were excised from the chop using a cork borer (25 mm diameter) and weighed. Later, the cores of muscle were placed into a DL plastic container (Christensen Aps Industrivaengetand, Hilleroed, Denmark) and stored at 4°C. Forty-eight h after sampling, the sample was removed from the container, carefully dabbed and weighed. DL was finally estimated by calculating the difference between the initial and the final weight of the muscle sample.

### Statistical Analysis

All statistical analyses were performed using the SAS 9.1 software (SAS Institute, Inc. 2002), with each group of pigs as an experimental unit. Behavioural data recorded during unloading and handling towards the restrainer were analysed with a Wilcoxon-Mann-Whitney test. In lairage, behavioural data were analysed with a Friedman test for continuous measures (drinking and aggressive behaviours), and by analysis of variance with repeated measures for postures (lying down or standing). Corticoids and catecholamines were analysed after log transformation to correct for deviation from normality, which was assessed by a Shapiro-Wilk test. These data were submitted to an analysis of variance according to a 2 × 2 factorial design in complete randomized blocks. Behavioural and hormonal data were only analysed for nine blocks, because observation conditions were different on the first week due to technical problems. Stomach data were also submitted to analysis of

variance, with the exception of content type data, which were analysed by a Chi-square test. An ANOVA was used to analyse the meat quality data according to a 2 × 2 factorial design in complete randomized blocks. For carcass contamination level, all bacterial counts were transformed to Log values and a Shapiro-Wilk test for normal distribution was applied. Log values of the means (log A) for sets of counts were calculated from the formula  $\text{Log } A = \bar{x} + \text{Log}_{10}(\text{SD}^2/2)$  (Brown and Baird-Parker 1982). The Log value of the lowest detection level ( $\text{Log}_{10}(5) = 0.69897$ ) was assumed for the calculation of  $\bar{x}$  and SD values for coliforms and *E. coli* counts when no colony was detected on the grid. A Log value for the total number of bacteria recovered (N) was calculated for each set of counts by adding the counts in each set and calculating the Log of the sum. These Log values were submitted to an analysis of variance with the option LIFEREG of SAS. The carcass trimmings taken on the dressing line were analysed with a Fischer test.

No interactions were found between handling quality and group size during lairage for any of the variables under study. Thus data were pooled across treatments and discussed according to the main effects of handling type and group size only.

## RESULTS

### Behavioural Observations

During unloading from the truck, rough handling increased the frequency of climbing ( $P < 0.05$ ), slipping ( $P < 0.01$ ) and turning around ( $P < 0.001$ ) (Table 1). In the stunning chute, rough handling only increased the frequency of climbing ( $P < 0.05$ ) (Table 1).

In lairage, the pigs kept in large groups were observed more often standing ( $P < 0.05$ ) than pigs in small groups, which spent more time lying ( $P < 0.05$ ). No significant differences in postures were observed between RH and GH pigs (Standing: RH = 55.8% ± 8.2 and GH = 55.7% ± 8.6 and lying down: RH = 44.2% ± 8.2 and GH = 44.3% ± 8.6). RH pigs were observed drinking less frequently than GH pigs (10.4% ± 5.3 vs. 19.1% ± 9;  $P < 0.01$ ), whereas no difference between groups sizes was recorded (group of 30: 13.6% ± 5.0 and group of 10: 16% ± 8.2). The pigs kept in large groups fought 10 times more than pigs in small groups ( $P < 0.001$ ) and were more often observed in agonistic interactions ( $P < 0.05$ ) (Fig. 2). The type of handling used to drive pigs to the pen did not influence the frequency of fighting and agonistic interactions during lairage. The frequencies of fighting were 6.4% ± 3.5 and 4.8% ± 2.1, and the frequencies of agonistic interactions were 6.2% ± 1.1 and 8.7% ± 2.3, for pigs handled roughly and gently, respectively. Overall, at the start of the lairage period, 70 to 90% of pigs were standing, but this proportion decreased to 10 to 30% at the end of the first hour (Fig. 3).

### Physiological Measures

The type of handling had no impact on any physiological stress indicators in the urine at slaughter, except for the cortisol concentration (Table 2). Indeed, cortisol level increased slightly ( $P < 0.05$ ), although to a little extent, in

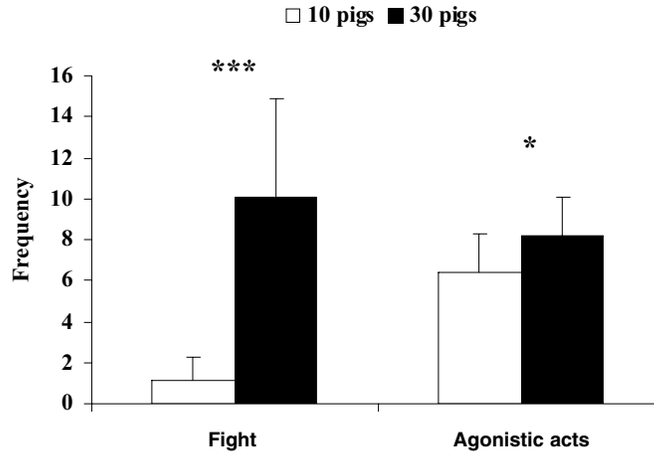
**Table 1. Effect of handling quality (rough vs. gentle) on behaviour of pigs during unloading and in the stunning raceway on pigs from groups of 10 (mean ± SD)<sup>a,y</sup>**

| Variables      | Unloading |           | P   | Stunning raceway |           | P  |
|----------------|-----------|-----------|-----|------------------|-----------|----|
|                | RH        | GH        |     | RH               | GH        |    |
| Climbing       | 0.6 ± 0.3 | 0         | *   | 3.8 ± 0.8        | 0.6 ± 0.4 | *  |
| Turning around | 5.1 ± 0.7 | 1.1 ± 0.4 | *** | 4.4 ± 0.9        | 5.0 ± 0.4 | NS |
| Slipping       | 1.3 ± 0.3 | 0.1 ± 0.1 | **  | –                | –         |    |

<sup>a</sup>RH = rough handling; GH = gentle handling.

<sup>y</sup>– = no measures.

\*, \*\*, \*\*\* P < 0.05, P < 0.01, and P < 0.001, respectively; NS = not significant.



**Fig. 2.** Frequency of fighting and agonistic acts (bites and head knocks) during lairage according to group size (10 pigs vs. 30 pigs; mean ± SD; \*P < 0.05; \*\*\*P < 0.001). Observation made by scan sampling on a subsample of 10 pigs within each group of 30.

the urine of pigs kept in small groups compared with those in larger groups.

**Stomach Weight and Content Composition**

The handling quality and group size during lairage did not affect stomach weight (full and empty) or DM of contents (Table 3). The number of stomachs containing liquid, sawdust or feed was not different between treatments either (Table 4). However, most of the stomachs evaluated in this study contained mainly liquid (76.5%). Only 17.7 and 5.8% of stomachs contained feed and sawdust, respectively. In addition, taking into account the criteria set by Chevillon (1994) to assess the fullness of stomachs due to incorrect fasting (≤ 1.4 kg for stomach weight and ≤ 500 g for content weight), 24.6% of the stomachs were full or partially full and of that portion 79.4% contained mainly liquid.

**Carcass Hygiene**

The handling quality (rough vs. gentle) and group size (10 vs. 30 animals) applied during lairage had no significant impact on the level of microbial carcass contamination whether it was for the TAM, coliforms or *E. coli* counts (Table 5). The 400 TAM counts were all well below 3 Log<sub>10</sub> CFU cm<sup>-2</sup>, and coliforms and generic *E. coli* were all below 60 Log<sub>10</sub> CFU cm<sup>-2</sup>. Furthermore, no coliforms or *E. coli* were detected on 15 and 30 of the 400 grids analysed, respectively. During evisceration, only one stomach was

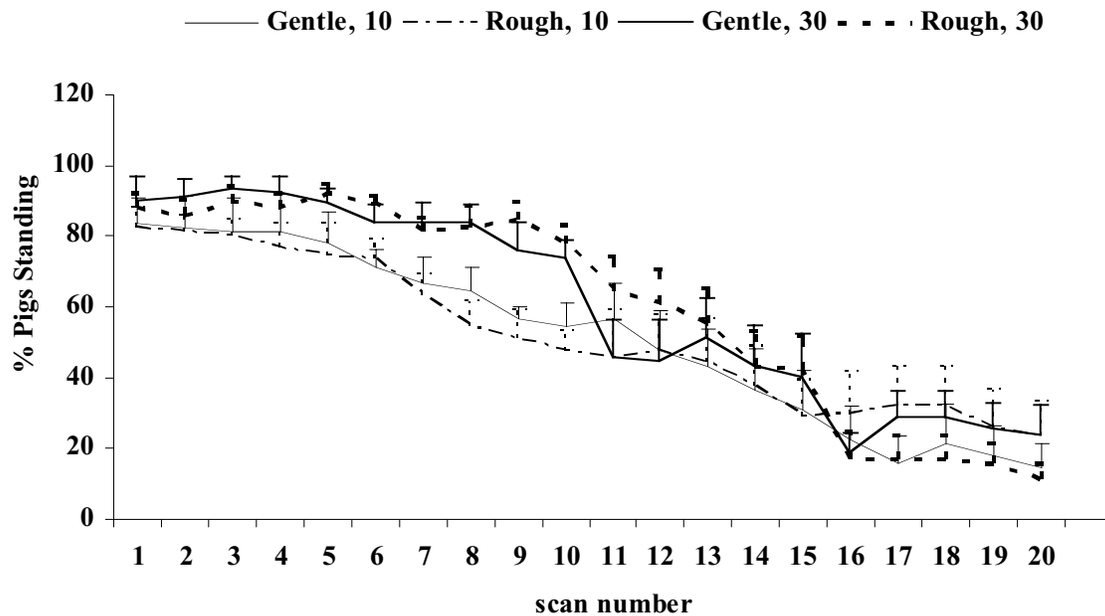
perforated. The number of pig carcasses per treatment from which meat was trimmed off by the veterinary inspectors and the abattoir employees on the dressing line because of visual contamination and defects was not different between treatments (GH = 32, RH = 42 and group of 10 pigs = 42, 30 pigs = 31).

**Skin Bruise and Meat Quality**

Bruise score tended to be higher (P = 0.06) in RH compared with GH pigs (Table 6). The type of handling also influenced pork quality as RH pigs had lower (P < 0.01) pH<sub>1</sub> and higher DL, EC and a\* (redness) values (P < 0.05, P < 0.01 and P < 0.05, respectively) in the LT muscle. Group size only affected the fall in pH 24 h post mortem with the pH<sub>u</sub> value being slightly higher (P < 0.01) in the LT muscle of the larger group size pigs.

**DISCUSSION**

There is evidence that a shock with an electric prod is more aversive to pigs than inhaling 90% CO<sub>2</sub> (Jongman et al. 2000) and can cause bruises on the carcass and lead to poor meat quality (Faucitano 2001a; Hemsworth et al. 2002). For this reason, the use of electric goads must be very limited (shocks lasting < 2 s; EC 1993). The results of this study revealed that the use of the electric prod as part of the rough treatment, although limited, produced panic within the group, increasing escape attempts at unloading and at the



**Fig. 3.** Percentage of pigs standing during lairage according to handling quality (gentle vs. rough) and group size (10 pigs vs. 30 pigs; mean  $\pm$  SD). Observation made by scan sampling (total of 20 scans) on a subsample of 10 pigs within each group of 30 pigs.

**Table 2.** Effect of handling quality (rough vs. gentle) and group size in pigs (10 vs. 30 per group) on urinary hormone levels at slaughter

| Variables                                      | Handling <sup>z</sup> |       |      | <i>P</i> | Group size |                 |      | <i>P</i> |
|--|-----------------------|-------|------|----------|------------|-----------------|------|----------|
|  | RH                    | GH    | SEM  |          | 10         | 30 <sup>y</sup> | SEM  |          |
| Cortisol (ng mg <sup>-1</sup> creatinine)      | 14.82                 | 13.29 | 0.79 | NS       | 14.94      | 13.12           | 0.79 | *        |
| Cortisone (ng mg <sup>-1</sup> creatinine)     | 23.66                 | 21.66 | 1.27 | NS       | 23.29      | 21.49           | 1.23 | NS       |
| Noradrenaline (ng mg <sup>-1</sup> creatinine) | 16.02                 | 15.52 | 0.58 | NS       | 14.76      | 16.84           | 0.65 | NS       |
| Adrenaline (ng mg <sup>-1</sup> creatinine)    | 8.26                  | 8.11  | 0.35 | NS       | 7.70       | 8.70            | 0.36 | NS       |
| Dopamine (ng mg <sup>-1</sup> creatinine)      | 13.17                 | 12.83 | 0.59 | NS       | 13.14      | 12.86           | 0.63 | NS       |

<sup>z</sup>RH = rough handling, GH = gentle handling.

<sup>y</sup>Observation made on a subsample of 10 pigs within each group of 30.

\**P* < 0.05; NS = not significant.

**Table 3.** Effect of handling quality (rough vs. gentle) and group size in pigs (10 vs. 30 per group) on stomach weights at slaughter

| Variables           | Handling <sup>z</sup> |       |      | <i>P</i> | Group size |                 |      | <i>P</i> |
|---------------------|-----------------------|-------|------|----------|------------|-----------------|------|----------|
|                     | RH                    | GH    | SEM  |          | 10         | 30 <sup>y</sup> | SEM  |          |
| Full stomach (g)    | 817.4                 | 810.5 | 18.3 | NS       | 797.4      | 830.5           | 19.7 | NS       |
| Emptied stomach (g) | 413.1                 | 413.3 | 4.1  | NS       | 412.1      | 414.2           | 4.2  | NS       |
| Stomach content (g) | 381.7                 | 376.1 | 15.6 | NS       | 363.4      | 394.5           | 16.9 | NS       |
| DM (%) <sup>x</sup> | 14.5                  | 13.9  | 0.4  | NS       | 14.3       | 14.0            | 0.4  | NS       |

<sup>z</sup>RH = rough handling; GH = gentle handling.

<sup>y</sup>Observation made on a subsample of 10 pigs within each group of 30.

<sup>x</sup>DM = dry matter.

NS = not significant.

entrance into the stunning chute, thus making the group more difficult to handle.

The quality of handling from the unloading deck to the pen only had an effect on drinking behaviour during lairage. GH pigs drank two times more frequently than RH pigs. Considering that GH pigs did not receive electric shocks during handling, it is possible that they were less stressed and adapted faster to their new environment than RH pigs.

This result confirms the observation of Warriss (2000) regarding the effect of previous handling on the behaviour of pigs in the lairage pen.

In practice, increased electric goading is often observed at the entrance into the stunning chute as the handler has no other means of separating a group of pigs and driving them individually down the stun race at a constant speed. In this study, climbing behaviour was observed more often than

**Table 4. Effect of handling quality (rough vs. gentle) and group size in pigs (10 vs. 30 per group) on number of stomachs containing liquid, sawdust or feed**

| Variables | Handling <sup>z</sup> |     | Group size |                 |
|-----------|-----------------------|-----|------------|-----------------|
|           | RH                    | GH  | 10         | 30 <sup>y</sup> |
| Liquid    | 142                   | 148 | 150        | 140             |
| Feed      | 35                    | 32  | 29         | 38              |
| Sawdust   | 8                     | 14  | 7          | 15              |

<sup>z</sup>RH = rough handling; GH = gentle handling.

<sup>y</sup>Observation made on a subsample of 10 pigs within each group of 30.

turning around at this stage, as a result of the combined effect of electric goading and reduced space allowance. It is well documented that when space is decreased, turning around is limited and pigs try to escape from the stressor (the electrical shock and the handler in this study) by climbing over the backs of other pen mates in search of protection within the group (Guise and Penny 1989; Lambooj and Engel 1991).

In our study, 70 to 90% of the pigs laid down approximately 1 h after they were in lairage. A similar finding was reported by Moss (1978) and Geverink et al. (1996), and confirms the benefit of providing pigs with a period of rest to allow them to recover from transport and the associated handling (Pérez et al. 2002; Warriss 2003). The increased level of activity and aggressiveness reported in pigs kept in the larger group is consistent with the findings by Bryant and Ewbank (1972), Petherick (1983) and Geverink et al. (1996), but contrasts with those of Andersen et al. (2004). According to Turner et al. (1999), the behaviour of pigs kept in large groups compared with small groups depends on their perception of the group size and on the availability of water and food in the pen. If the space per pig and resources (food, water, etc.) are sufficient, it appears that the level of activity and the frequency of agonistic acts do not increase. As all pigs were fasting, the aggressiveness observed in this study, especially in the large groups, might have been increased by the lack of food. Indeed, fasting often results in increased fighting, especially in mixed pigs (Fernandez et al. 1995; Brown et al. 1999; Turgeon 2003).

Rough preslaughter handling and poor social environment in lairage may cause psychological and physical stress in pigs as shown by the significant increase in blood hormone levels, heart rate and body temperature reported in several studies (Faucitano 2001b; Hemsworth et al. 2002). However, except for a slight effect of group size on cortisol, the treatments applied had no effect on the concentrations of urinary hormones at slaughter in this study. Overall, the urinary concentrations of CA recorded in this study were higher than those reported by Mormède et al. (2004) and lower than those reported by Foury et al. (2005) for pigs slaughtered in a commercial abattoir following transportation. On the other hand, cortisol concentrations were lower than those reported by Mormède et al. (2004) and Foury et al. (2005). Discrepancies between studies may be due to genetic differences (Hay and Mormède 1998; Mormède et al. 2004). Overall, the lack of strong differences between treatments suggests that the pre-slaughter management applied

in this study was stressful for all pigs and may have masked treatment effects. For instance, the lairage environment was noisy (82 to 108 dB). Geverink et al. (1998) showed that the noise produced by the machinery, pressure hoses, and pig and human vocalisations represented a source of stress which evoked huddling and escape behaviour. Talling et al. (1996) and Kanitz and Tuchscherer (2005) also reported an increased blood cortisol concentrations and heart rate in pigs exposed to high sound intensity levels (85–97 dB).

Although treatments affected the behaviour of pigs during handling and lairage, they had no impact on stomach weight. This contrasts with the findings by Enck et al. (1989) who reported gastric emptying was delayed in stressed rats. However, it is consistent with the physiological response of pigs, which suggests that they were equally stressed, regardless of treatment. In the present study, despite fasting for 20 h, 23.5% of stomachs were not empty according to the criteria of Chevillon (1994). Of the stomachs that did not meet the criteria, a high percentage (79.4%) contained liquid. However, it should be noted that treatments had no impact on stomach content, despite the fact that GH pigs drank more than RH pigs. The reason for the lack of impact of drinking behaviour during lairage on stomach content weight and composition can be twofold. On one hand, RH pigs might have drunk more during the last 2 h of lairage (not under observation), and compensated for their lower drinking frequency in the first hour. On the other hand, given that liquid evacuation rate from the stomach is higher than that of solid content (Gregory et al. 1990), the water consumed by pigs during the first hour of lairage may have been absorbed or evacuated before slaughter.

Given the precautions taken during the removal of the gastro-intestinal tract from the carcass to avoid content spillage at sampling, only one stomach was perforated during evisceration. Hence, the treatments applied in this study did not appear to affect the efficacy of the evisceration procedure. The lack of difference between treatments in microbial carcass contamination and in the number of carcasses trimmed for visual contamination and defects suggest that the stress conditions applied in this study had limited impact on the carcass microbial quality. Overall carcass contamination was very low in our experiment. The levels of TAM, coliforms or *E. coli* counts measured on all carcasses were within the guidelines of the Meat Hygiene Manual [Canadian Food Inspection Agency (CFIA) 2006], independent of the treatment applied. Carcass trimming occurred mainly on week 10 (16.3% of the carcasses), for no obvious reason. This suggests that carcass contamination at that time may have been caused by an unexpected event independent of the preslaughter handling conditions under study. Botteldoorn et al. (2003) showed a high variation in the incidence of contaminated carcasses between different sampling days both at the same abattoir (between 3 and 52%) and between abattoirs (between 0 and 70%).

The increased climbing behaviour provoked by electric prodding of pigs at unloading and right before slaughter increased the incidence of skin bruises on the carcass. Bruises may have been produced by the fore nails of pigs climbing onto the back of others while escaping from the

**Table 5. Effect of handling quality (rough vs. gentle) and group size in pigs (10 vs. 30 per group) on carcass contamination: TAM (total aerobic mesophilic), coliforms and *E. coli* counts**

|                |                       |                 | Statistics  |                 |                    |                |                |
|----------------|-----------------------|-----------------|-------------|-----------------|--------------------|----------------|----------------|
|                |                       |                 | $\bar{x}^x$ | SD <sup>w</sup> | Log A <sup>v</sup> | N <sup>u</sup> | % <sup>t</sup> |
| TAM            | Handling <sup>z</sup> | RH              | 1.73        | 0.32            | 1.85               | 4.16           | 100            |
|                |                       | GH              | 1.69        | 0.32            | 1.81               | 4.10           | 100            |
|                | Group size            | 10              | 1.73        | 0.33            | 1.85               | 4.16           | 100            |
|                |                       | 30 <sup>y</sup> | 1.69        | 0.32            | 1.81               | 4.11           | 100            |
| Coliforms      | Handling              | RH              | 1.68        | 0.69            | 2.23               | 5.20           | 79             |
|                |                       | GH              | 1.71        | 0.64            | 2.18               | 4.96           | 78             |
|                | Group size            | 10              | 1.73        | 0.67            | 2.24               | 5.19           | 82             |
|                |                       | 30 <sup>y</sup> | 1.66        | 0.66            | 2.16               | 4.97           | 75             |
| <i>E. coli</i> | Handling              | RH              | 1.45        | 0.69            | 1.99               | 5.09           | 52             |
|                |                       | GH              | 1.49        | 0.57            | 1.86               | 4.34           | 68             |
|                | Group size            | 10              | 1.48        | 0.58            | 1.87               | 4.77           | 65             |
|                |                       | 30 <sup>y</sup> | 1.46        | 0.67            | 1.98               | 4.94           | 55             |

<sup>z</sup>RH = rough handling; GH = gentle handling.

<sup>y</sup>Observation made on a subsample of 10 pigs within each group of 30.

<sup>x</sup>Mean of Log<sub>10</sub> CFU cm<sup>-2</sup> for TAM, mean of Log<sub>10</sub> CFU 983 cm<sup>-2</sup> for coliforms and *E. coli* counts.

<sup>w</sup>Standard deviation of the log<sub>10</sub> counts.

<sup>v</sup>Log<sub>10</sub> of the arithmetic mean.

<sup>u</sup>Log<sub>10</sub> of the total number recovered from 10 × 983 cm<sup>2</sup>.

<sup>t</sup>Percentage of samples with countable colonies on the grids.

**Table 6. Effect of handling quality (rough vs. gentle) and group size in pigs (10 vs. 30 per group) on meat quality traits**

| Variables                            | Handling <sup>z</sup> |       | SEM  | <i>P</i> | Group size |                 |      | <i>P</i> |
|--------------------------------------|-----------------------|-------|------|----------|------------|-----------------|------|----------|
|                                      | RH                    | GH    |      |          | 10         | 30 <sup>y</sup> | SEM  |          |
| Skin bruise score <sup>x</sup>       | 1.53                  | 1.36  | 0.06 | 0.06     | 1.46       | 1.43            | 0.06 | NS       |
| pH <sub>1</sub>                      | 6.01                  | 6.11  | 0.05 | **       | 6.05       | 6.06            | 0.05 | NS       |
| pHu                                  | 5.60                  | 5.58  | 0.02 | NS       | 5.58       | 5.61            | 0.02 | **       |
| EC <sup>w</sup> (μS)                 | 5.84                  | 4.97  | 0.22 | **       | 5.45       | 5.36            | 0.22 | NS       |
| Subjective colour score <sup>y</sup> | 2.45                  | 2.42  | 0.05 | NS       | 2.42       | 2.44            | 0.05 | NS       |
| <i>L</i> <sup>*</sup>                | 52.44                 | 52.17 | 0.28 | NS       | 52.26      | 52.36           | 0.28 | NS       |
| <i>a</i> <sup>*</sup>                | 8.72                  | 8.44  | 0.13 | *        | 8.64       | 8.52            | 0.13 | NS       |
| <i>b</i> <sup>*</sup>                | 6.47                  | 6.34  | 0.15 | NS       | 6.45       | 6.35            | 0.15 | NS       |
| Drip loss (%)                        | 7.58                  | 6.96  | 0.29 | *        | 7.38       | 7.15            | 0.29 | NS       |

<sup>z</sup>RH = rough handling; GH = gentle handling.

<sup>y</sup>Observation made on a subsample of 10 pigs within each group of 30.

<sup>x</sup>According to photographic standards (from 1 = none to 5 = severe; MLC 1985).

<sup>w</sup>EC = electrical conductivity measured by Pork Quality Meter (Intek, Aichach, Germany; > 5 μS = fast post mortem glycolysis leading to unacceptable pork; Barton-Gade et al. 1996)

<sup>y</sup>According to Japanese Color Scales (from 1 = pale to 6 = dark; Nakai et al. 1975).

\*, \*\* *P* < 0.05 and *P* < 0.01, respectively; NS = not significant.

source of stress (handler and electric goad) in a situation of high stocking density. This has been previously reported by Faucitano et al. (1998) in pigs lined up and handled with electric prods in a stunning chute not equipped with hold-down bars to prevent climbing activity.

The direct effect of electric prodding on the production of exudative pork is extensively reported in the literature (D'Souza et al. 1998; Støier et al. 2001; Hemsworth et al. 2002; Hambrecht et al. 2005). Van der Wal et al. (1999) reported that only a very short-term stress (1 min) immediately before stunning can be sufficient to produce exudative pork. Likewise, Hemsworth et al. (2002) reported that the nature of the interaction received by the pigs (pats and strokes or slaps and electric goad hits) in the crowding pen prior to the final race leading to the stunning area was direct-

ly correlated with the fear of the handler. In the present study, the increased physical activity (climbing) a few minutes before slaughter caused by the presence of the handler in combination with the electric shock generated by the prod reduced muscle pH<sub>1</sub> and water-holding capacity of pork meat.

Despite the higher levels of activity and aggressiveness observed in the pigs kept in large groups, group size had no effect on skin bruise score and had just a small, but not biologically significant, effect on pH<sub>u</sub>. These results are surprising as fighting in lairage is usually associated with high bruise scores on the carcass and high muscle pH<sub>u</sub> due to the effects of physical activity on glycogen levels at slaughter (Warriss 1996; Gispert et al. 2000). Nevertheless, our results may support the findings by Andersen et al. (2004), who

reported that the proportion of pigs not participating in fights was higher in a large group. According to Pagel and Dawkins (1997), the probability of observing the same individuals fighting declines sharply with increasing group size, because the energetic costs and physical injury associated with establishing and maintaining a social hierarchy in a large group may encourage the adoption of an alternative strategy. Thus, the fact that fewer animals may have been involved in fights in the large group, might explain the lack of effect of increased agonistic acts on the average skin bruise score and pork quality of the group.

### CONCLUSIONS

Poor handling increased the incidence of flight behaviours at unloading and aggressive behaviours along the lairage alleys and resulted in poorer pork quality. The level of activity and the frequency of aggression during lairage also increased with group size. However, group size had no impact on pork quality. The stressors applied on pigs in lairage did not influence stomach weight at slaughter, suggesting that they did not contribute to stomach emptying variation before slaughter. Furthermore, all samples (400 in total) were within the guidelines of the Meat Hygiene Manual for the indicator organisms of the carcass hygienic status (TAM, coliforms or *E. coli* counts), independent of the treatments applied (CFIA 2006). Hence, no treatment tested had a negative effect on the hygiene level observed and all carcasses respected the food safety standards for the commercial sale. Further research is needed to study the effect of drinking rate during lairage on stomach weight and content composition.

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