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# Purified natural pig immunoglobulins can substitute dietary zinc in reducing piglet post weaning diarrhoea

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- 4 Short communication
- 5 Purified natural pig immunoglobulins can substitute dietary zinc in
- 6 reducing piglet post weaning diarrhoea
- 7
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- 16

17 Abstract

18 Enteric infectious disease in weaner piglets, including postweaning diarrhoea (PWD), are

19 usually treated and/or prevented with antibiotics and/or zinc oxide in the piglet feed.

20 However extensive use of antibiotics and zinc oxide in intensive animal production is 21 unwanted as it may promote microbial antibiotic resistance and pose environmental 22 problems. Recently, in an experimental model of PWD, we observed that oral 23 administration of purified porcine immunoglobulin G (pplgG) from pooled natural pig 24 plasma could reduce enteric infection. In the present study we were able to reproduce 25 these results as it was observed that oral pplgG accelerated clearance of faecal 26 haemolytic bacteria in pigs challenged with E. coli in comparison with pigs not receiving 27 pplgG. This effect was observed upon feeding pplgG for seven days postweaning 28 suggesting that ppIgG does not have to be used prophylactically for several days 29 preweaning. Furthermore, the effect of oral administration of pplgG for seven days 30 postweaning was equal to or better than that of dietary zinc oxide in reducing diarrhoea 31 symptoms and in clearing faecal haemolytic bacteria for 14 days postweaning. These 32 observations warrant future trials of dietary pplgG in intensive swine production units to 33 establish its performance as an alternative to dietary antibiotics and zinc oxide for 34 preventing PWD.

#### 35 Keywords

Postweaning diarrhoea; Immunoglobulins; IgG; Zinc oxide; Antibiotics; ETEC; E. coli F4;
Feed supplement; Feacal score

38

#### 39 1. Introduction

40 After parturition, lactogenic immunity provides offspring with oro-gastric protection 41 against infectious pathogens (Hedegaard and Heegaard, 2016). In the conventional 42 swine production systems piglets are weaned at an immunologically immature age (3–4 43 weeks) depriving them of the continued supply of protective lactogenic antibodies at the 44 same time as they are placed in a new environment with increased risk of enteric 45 bacterial infections that may lead to postweaning diarrhoea (PWD) (Madec et al., 1998). 46 PWD is characterised by diarrhoea caused by enteric infection by Enterotoxigenic 47 Escherichia coli (ETEC) usually within three days postweaning (Fairbrother et al., 2005). 48 PWD, like other enteric infections in pigs, can be treated with dietary antibiotics 49 (primarily tetracyclines, penicillins and macrolides (Becker, 2010; DANMAP, 2015; 50 Wageningen University, 2012)), and/or zinc oxide (Pluske, 2013). Indeed, antibiotics and 51 zinc can improve average daily growth (ADG) in weaner piglets (Cromwell, 2002; Molist 52 et al., 2011), and dietary zinc can reduce frequency of PWD in weaner piglets ( Owusu-53 Asiedu et al., 2003 ; Pluske, 2013). The mechanisms by which zinc oxide increases ADG 54 and reduces diarrhoea in weaner piglets are not fully understood but seem to involve 55 both restoring plasma zinc levels to normal in piglets after weaning (Davin et al., 2013) 56 and improving intestinal homeostasis (Liu et al., 2014a; Liu et al., 2014b ; Shen et al., 2014). 57

However, the widespread use of dietary zinc oxide in intensive animal production can result in pollution of farmlands and groundwater through repeated fertilisation with zinc-containing residual manure (Hill et al., 2005). In addition, zinc oxide in combination with antibiotics appears to accelerate microbial antibiotic resistance by increasing the rate of the exchange of antibiotic-resistance-gene containing plasmids in the microbiota community in both soil and piglet intestines (Lin et al., 2016; Pal et al., 2015 ; Vahjen et al., 2015); thus, there is a need to reduce both kinds of interventions.

As a sustainable and economically feasible alternative to antibiotics and zinc for treating
PWD, we have previously investigated the use of natural purified porcine

67 immunoglobulin G (pplgG) from pooled abattoir blood plasma (Hedegaard et al., 2016). First, it was established that natural pplgG does indeed contain immunoglobulin 68 69 dependent anti-E. coli activity. Secondly it was shown that a dietary pplgG supplement 70 in a model of PWD led to faster clearance of an ETEC challenge infection than seen in a 71 comparable control group not provided with dietary pplgG (Hedegaard et al., 2016). 72 This prompted us to further investigate this effect of ppIgG in E.coli challenge models. A 73 long term, low pplgG dose experiment and a short term, high pplgG dose experiment 74 were performed. In both experiments dietary pplgG resulted in reduction of diarrhoea 75 and in number of faecal haemolytic bacteria; moreover, in the second trial we observed 76 that pplgG reduced diarrhoea and cleared the enteric infection faster than dietary zinc 77 oxide. These observations warrant future experiments investigating the use of dietary 78 ppIgG postweaning as an alternative to dietary zinc oxide and antibiotics in 79 treating/preventing PWD.

80

#### 81 2. Materials & methods

The study comprised two infection experiments, both using E. coli O149 challenge at two consecutive days post weaning. In experiment 1 (long term, low IgG dose pilot study) 750 mg/day of ppIgG was provided orally for five days before and 10 days post weaning. In experiment 2 (short term, high IgG dose) 1.9 g of ppIgG was given orally twice daily for seven days after weaning/challenge. Experiment 2 also comprised a group receiving dietary zinc oxide for 10 days after weaning.

88

89 2.1. Purified porcine immunoglobulin G (ppIgG)

The ppIgG was prepared from pooled pig plasma by expanded bed chromatography (EBA) at Upfront Chromatography A/S (Copenhagen), as described previously (Hedegaard et al., 2016). Concentrated porcine blood plasma was obtained from Daka SARVAL A/S (Lunderskov, Denmark). The batch of ppIgG used in first experiment (experiment 1, see below) was the same as in (Hedegaard et al., 2016), whereas a new batch of ppIgG was prepared prior to the second (experiment 2, see below).

96

97 2.2. ELISA

98 The IgG concentration in the batches of ppIgG was measured by a sandwich ELISA 99 (Hedegaard et al., manuscript in preparation), utilising a goat anti-pig IgG (Fc) antibody 100 (AAI41, Nordic Biosite ApS, Copenhagen) both for capture and detection. The IgG-101 concentration of ppIgG used in experiment 1 was 37.5 mg/ml and in experiment 2 the 102 ppIgG concentration was 75 mg/ml.

103 Anti-E. coli activity was found in both batches of pplgG, used in this study, by indirect 104 whole-E. coli cell ELISA previously reported (Hedegaard et al., 2016); briefly 96 wells flat 105 bottom microtiter plates (Maxisorp, NUNC, Thermo Scientific, Denmark) were coated 106 with 100  $\mu$ l fixed E. coli O138 (in-house strain isolated from piglet with diarrhoea) in 0.1 107 M sodium carbonate buffer pH 9.6 (OD546 = 0.25) at 4 °C overnight. All subsequent 108 operations were performed at room temperature. Next day wells were washed four 109 times in PBS with 0.05% Tween 20 (PBS-T), and blocked with 200  $\mu$ l PBS-T with 1% 110 bovine serum albumin (BSA; Sigma-Aldrich, Brøndby, Denmark) for 30 min when shaking 111 then followed by four times wash as above. The ppIgG was added in 2-fold dilution 112 series (diluted in PBS-T 1% BSA from 10 to 0.02 mg/ml). After 1 h of incubation with 113 shaking and 4 washes in PBS-T, detection antibody HRP-conjugated goat anti-pig IgG 114 (GGHL-5P; ICL, SMS Gruppen; Rungsted, Denmark) diluted 1/2000 in PBS-T 1% BSA was 115 added and incubated for 1 h with shaking. After washing, plates were developed by TMB 116 Plus substrate (Kem-En-Tec, Taastrup, Denmark), 100  $\mu$ l/well, and stopping colour 117 development by 100  $\mu$ l/well 0.5 M H2SO4 (VWR—Bie & Berntsen A/S). Optical density 118 was measured by a Thermo Scientific Multiscan EX microplate reader at 450 nm 119 subtracting background absorbance at 650 nm.

Using the antigen specific ELISA it was found that the pplgG batch used in experiment 1
had lost 14% of activity in comparison to the original plasma pool, whereas the batch
used in experiment 2 had lost no activity (data not shown).

123

124 2.3. Experiments

125 2.3.1. Experimental procedure

126 Two separate factorial experiments at Aarhus University, Foulum, involving 12 (10.8 ± 127 1.8 kg BW, from 2 litters) and 18 (7.3  $\pm$  1.0 kg BW, from 2 litters) pigs for the first and 128 second experiments, respectively, were conducted (see Table 1). In both experiments, 129 piglets were weaned from sows that had been tested to be homozygote carriers of the 130 dominant gene encoding for intestinal F4 fimbriae receptors (Van Haeringen 131 Laboratorium, b.v., Wageningen, The Netherlands) on DNA extracted from hair sample. 132 Piglets were weaned from the sows at day 28 of age, and were challenged with E. coli 133 O149:F4 on two consecutive days (d 29 and 30 of age). Within each experiment, piglets 134 from different litters were equally distributed among treatments.

In the first experiment, half of the piglets (Table 1; Exp. 1, Group 1 + 2) received once
daily for 15 days (5 days preweaning and 10 days postweaning) 20 ml (750 mg) of pplgG,

provided by a 20 ml syringe; a small plastic tube was connected to the syringe and the
pplgG was slowly dipped in and piglets willingly lapped up, ensuring that no pplgG was
lost.

The other half of the pigs received 20 ml 0.9% NaCl (Table 1; Exp. 1, Group 3 + 4). Pigs
were provided the immunoglobulin product before feeding.

142 In the second experiment, pigs were allotted into three challenge-groups (Table 1; lower 143 part): two groups received no immunoglobulins but piglets were provided with 25 ml of 144 0.9% NaCl and provision of feed (from day of weaning) based on wheat, barley and 145 dehulled soybean, and with either 2500 ppm zinc oxide (Hammershøj Pharmacy, 146 Hammershøj, Denmark) for 14 days postweaning (Table 1; Exp. 2, Group 1), or no zinc 147 oxide (Table 1; Exp. 2, Group 2). The third group received ppIgG via 20 ml syringe on the 148 morning before weaning (day 27), and on the morning of weaning (day 28). The pplgG 149 feeding was continued twice daily by drench gun for 7 days postweaning with a 150 provision of 25 ml (1.9 g) of ppIgG twice daily (Table 1; Exp. 2, Group 3).

151

The animal experiment was conducted according to the personal license (Charlotte Lauridsen, J. nr. 2012-15-2934-00125) obtained by the Danish Animal Experiments Inspectorate, Ministry of Food, Agriculture and Fisheries, Danish Veterinary and Food Administration, and animals were followed by proper veterinary surveillance throughout the experiment.

157

158 2.3.2. Animals, feeding and housing

159 All piglets were tested susceptible to E. coli O149:F4 by using a DNA marker genotyping-160 based test (van Haeringen laboratorium b.v., Wageningen, The Netherlands) on DNA 161 extracted from hair samples. From weaning and onwards, the pigs had ad libitum access 162 to feed and water. The feed was a standard diet for weaners prepared at the facilities at 163 Aarhus University, Foulum. Two littermates of similar weight were housed in pairs in 164  $1.45 \text{ m} \times 1.7 \text{ m}$  pens with concrete flooring and sawdust bedding. In experiment 1, the 165 unchallenged (Groups 1 + 3) and challenged pigs (Groups 2 + 4) were housed in different 166 rooms to avoid cross contamination with the challenge E. coli strain. A door separated 167 the rooms between the challenged and unchallenged piglets, and personnel changed 168 clothes/shoes before entering any of the rooms. Environmental conditions including 169 temperature (~24 °C), humidity (~50%) and bedding (sawdust) were similar in the two 170 rooms. Experiment 2 was conducted in the same rooms, but all pigs were challenged 171 with E. coli, and there was free passage between the rooms throughout the entire 172 experiment.

173

#### 174 2.3.2.1. E. coli challenge

175 In both experiments, pigs were orally inoculated with 1.0 × 109 colony-forming units 176 (cfu) of E. coli O149:F4 in 5 ml 0.9% NaCl on day 1 and 2 after weaning (day 0 was the 177 day of weaning) using a syringe. After inoculation, the tube was flushed with 178 approximately 10 ml 10% NaHCO3 in order to neutralize gastric acid and increase the 179 survival rate of the challenge strain in the stomach, and to ascertain that all the E. coli 180 suspension had been given to the piglets. The control pigs (exp. 1) received equivalent 181 amounts (approximately 5 ml) of 0.9% NaCl and 10 ml 10% NaHCO3 in order to obtain an equal level of stress associated with the oral inoculation as for the challengetreatment.

184

185 2.3.3. Performance and diarrhoea assessment

186 Diarrhoea assessment was based on the consistency of the faeces (1 = hard, dry and 187 cloddy, 2 = firm, 3 = soft with shape, 4 = soft and liquid, 5 = watery and dark, 6 = watery 188 and yellow, 7 = foamy and yellow) from the day prior to challenge until 7 days after. A 189 faecal consistency score >3 was defined as a clinical sign of diarrhoea (Carstensen et al., 190 2005). Before the E. coli challenge, and on day 2 after challenge, and daily until day 5 191 after challenge, and thereafter every second day during the second week after weaning, 192 faecal samples were collected from the rectum of the pigs and 1 g faeces was suspended 193 in a (1:10, wt/wt) peptone solution and homogenized by bag mixer (BagMixer100, 194 Interscience, St. Nom, France). Serial dilutions of the slurry were done prior to enumeration of haemolytic E. coli on blood agar (BA; Oxoid) after aerobic incubation at 195 196 37 °C overnight. From each BA plate, five haemolytic E. coli colonies were selected and 197 tested for O149 and O138 type reactions by O-seroagglutination (Statens Serum Institut, 198 Copenhagen, Denmark). In addition, faecal samples were analysed for dry matter by 199 freeze-drying (ScanVac Coolsafe 55, Labogene Aps, Lynge, Denmark).

Feed intake was recorded daily for each pen and body weight of the pigs was recorded at the beginning and weekly thereafter until the end of the experiment. Average daily feed intake (ADFI) and gain (ADG) were determined based on pen by dividing the total feed intake or total weight gain of pigs in each pen by days of feeding.

205 2.4. Statistics

The effects of pplgG, dietary zinc oxide or no treatment for seven days postweaning on diarrhoea symptoms and bacterial count in both experiments were statistically analysed using either Mann-Whitney test or Two-way ANOVA followed by Tukey's post-test, in GraphPad Prism version 7.00 for Windows, GraphPad Software, La Jolla California USA, www.graphpad.com.

211

#### 212 3. Results and discussion

In this study we performed two infection experiments using an E. coli O149 challenge strain at both day one and two after weaning. The administration of pplgG was different in the two experiments: In experiment 1, 750 mg of pplgG was provided orally once daily prophylactically for five days preweaning and then 10 days postweaning, whereas in experiment 2, 1.9 g of pplgG was administered orally twice daily for seven days postweaning. Two different pplgG batches were used for the two experiments, for experiment 1 a 37.5 mg/ml batch and for experiment 2 a 75 mg/ml batch.

220

#### 221 3.1. Disease

In both experiments diarrhoea was observed within one day after inoculation due to the O149 ETEC challenge as observed by faecal scoring (Fig. 1A + B) and faecal dry matter measurement (Fig. 1C + D). We observed in both experiments that the +pplgG +challenge group (group 2 in exp. 1 and group 3 in exp. 2) had, at day five, significantly less diarrhoea as compared to the control challenge groups (group 4 in exp. 1 and group 227 2 in exp. 2) that were not provided with pplgG as observed by a lower faecal scoring and 228 a higher percentage of faecal dry matter (p < 0.03; Fig. 1, days 1–5). In experiment 1 the O149 ETEC challenge strain began to cease at day 7, however an unintended enteral 229 230 infection, starting around day 5–7 after weaning and lasting the remainder of the study 231 period was observed (Fig. 1A + C, days 5–21), coinciding with the appearance of an O138 232 ETEC strain in faeces (detected by PCR; data not shown). The presence of this 233 unintended infection somewhat obscured experiment 1, however the data from day 0 to 234 five were promising and prompted us to conduct a second experiment in which the daily 235 IgG dose was quadrupled but then only provided seven days post weaning. This batch of 236 ppIgG had an IgG concentration of 75 mg/ml, and was administered twice daily, 237 amounting to a daily dose of 3.8 g.

Experiment 2 proceeded without any unintended infection, and group 3 (+pplgG +challenge group) had significantly less diarrhoea than both group 1 and 2 (+zinc +challenge and no treatment groups) within the first week after weaning (p < 0.02 Twoway ANOVA; Fig. 1B + D); demonstrating that pplgG significantly reduced diarrhoea in the PWD model within the first week post weaning to the same level or lower than dietary zinc.

Moreover, our results indicate that it is not necessary to use pplgG prophylactically for several days preweaning as there was a clear effect of pplgG in experiment 2 on diarrhoea without the five days preweaning pplgG treatment applied in experiment 1 (Fig. 1). This also corroborates a previous study (Hedegaard et al., 2016) where pplgG was mixed into the feed that was only available from weaning and decreased diarrhoeagenic Enterobacteriaceae in comparison to a non-pplgG diet. Also, it has been shown (Foged et al., 1986) that monoclonal anti-E. coli F4 fimbriae antibodies administered at as well as after challenge, but not prophylactically, protected neonatal piglets from an otherwise lethal challenge with F4+ ETEC. On the other hand, it seems to be important to maintain a high administration frequency and an adequately high antibody dose for limiting diarrhoea.

255

#### 256 3.2. Microbiology

257 In both experiments pplgG helped clear the challenge strain within one week of 258 challenge (Fig. 2, days 1–7). However, as noted above, all groups in experiment 1 259 experienced an unintended O138 ETEC infection from day 4–9, as seen by an increase in 260 numbers of faecal haemolytic bacteria (Fig. 2A, days 4 + 7 + 9). Group 1 (+pplgG no 261 challenge) experienced this infection at day 4 while it was observed in group 2 (+pplgG 262 +challenge) on day 9. Thus, the daily administration of 750 mg of natural IgGs did not 263 provide protection against the unintended infection in neither of the two +ppIgG groups 264 (Fig. 2A, groups 1 + 2, days 4-21), even though in vitro data support that ppIgG binds to 265 O138 (and O149) ETEC and can inhibit their adhesion to intestinal epithelial cells in vitro 266 (Hedegaard et al., 2016). This might indicate that the unintended infection was 267 multifactorial, and/or that the daily dose of 750 mg natural ppIgG used in this 268 experiment was not adequate to prevent this type of infection. Data from doseresponse field trials are needed before any further conclusions can be made on this 269 270 matter.

In experiment 2, only infection with the challenge-strain was observed. Confirming previous results (Hedegaard et al., 2016), pplgG caused a faster clearance of diarrhoeagenic (haemolytic) bacteria in group 3 than was observed in the other two groups (Fig. 2B). Thus day 7 was the last day on which faecal haemolytic bacteria were 275 detected in group 3 (+pplgG), while these bacteria could still be detected on day 9 in 276 group 1 (+zinc group), and in group 2 (control) one piglet still had faecal haemolytic 277 bacteria on day 15 (Fig. 2B). Taken together, pplgG appears to intervene with the 278 colonization by the ETEC challenge strain shortening the period of infection significantly 279 (Fig. 2B, Day 5, p < 0.02) and was as efficient as dietary zinc in reducing the infection 280 with the ETEC challenge strain. Although this suggests that pplgG can shorten the 281 duration of diarrhoea in weaner piglets by decreasing the number of faecal 282 diarrhoeagenic haemolytic bacteria, this should ideally take place without a perturbing 283 the composition of the normal intestinal microbiota. In experiment 1, no change in the 284 faecal non-haemolytic bacteria were initially observed (Fig. 2C, days 1-5) however, as 285 the unintended infection emerged the number of faecal non-haemolytic bacteria began 286 decreasing (Fig. 2C, days 7-21). In group 2 (+pplgG +challenge) the faecal non-287 haemolytic bacteria actually were not recovered for 3 days at the height of the 288 unintended infection (Fig. 2C, days 11-14), which coincided with the termination of 289 administration of pplgG. However in experiment 2, no changes in the count of faecal 290 non-haemolytic bacteria in any of the three other groups were observed (Figs. 2D), 291 indicating that pplgG does not intervene with the intestinal non-haemolytic commensal 292 microbiota and that the microbiota changes observed in experiment 1 was probably due 293 to the unintended infection. These observations are supported by preliminary next 294 generation sequencing data on the faecal microbiota composition of piglets fed pplgG showing no change in non-haemolytic commensals (unpublished data). 295

Disease (diarrhoea) frequency, growth and feed conversion are primary end points for swine producers however the two experiments described here comprised low numbers of piglets (n = 12/18) and pens (n = 4/3) making it very difficult to analyse growth data statistically. Also, the data on growth and feed intake showed a very large pig-to-pig 300 variation. For example, in experiment 2 half of the piglets in group 3 (+pplgG), for 301 unknown reasons, became anorexic during the third week resulting in almost no weight gain for the group. Therefore it will be interesting to observe how pplgG 302 supplementation will influence disease resistance, growth and feed conversion in field 303 trials incorporating an adequate number of pigs to allow for appropriate statistical data 304 305 analysis to be performed. In spite of being a small preliminary study the results shown 306 here do however demonstrate the ability of dietary pplgG to clear an enteric ETEC 307 infection and thus pplgG could be used as an alternative to dietary zinc.

308

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## 454 Tables

## 455 Table 1: Study setup

Experiment 1	Group 1	Group 2	Group 3	Group 4
	+pplgG	+ppIgG+E.coli	Control	+E.coli
Number of piglets	3	3	3	3
<i>E. coli</i> F4+ challenge	No	Yes	No	Yes
750 mg ppIgG (20 ml)	Yes	Yes	No	No
20 ml 0.9% NaCl solution	No	No	Yes	Yes
Avg. weight at weaning	11.2±3.1	10.6±1.7	10.4±1.9	11.0±1.2
(kg)				
Experiment 2	Group 1	Group 2	Group 3	
	+ E.coli +Zn	+E.coli	+E.coli +pplgG	
Number of piglets	6	6	6 (5) <sup>1</sup>	
E. coli F4+ challenge	Yes	Yes	Yes	-
2x1.9 g pplgG (2x25 ml)	No	No	Yes	
25 ml 0.9% NaCl solution	Yes	Yes	No	
Zinc oxide in feed	Yes	No	No	
Avg. weight at weaning	6.9±1.1	7.9±0.9	7.0±0.9	
(kg)				

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<sup>1</sup> One piglet was euthanized on day 3 post infection due to serious illness.

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459 Figure legends

460 Figure 1. Faecal analysis: Faecal diarrhoea score (A+B) and faecal dry matter (C+D); (●) 461 no treatment, no E. coli; (●) no treatment, +E. coli: (-■-) +pplgG, no E. coli; (▲)pplgG 462 treatment, +E. coli; (=) dietary zinc, +E. coli. Each data point is plotted and curves 463 outline the mean for each group. Vertical dotted line indicates the last day of pplgG 464 administration. Pairwise comparisons between groups on each day were tested for 465 statistical significance using Mann-Whitney test: a = IgG vs. Zinc (p<0.03); b = IgG vs.Control (p<0.03); c = IgG vs. Zinc (p<0.02); d = IgG vs. Control (p<0.01); e = Zinc vs.466 467 Control (p<0.02).

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Figure 2. Faecal bacterial analysis: Content (CFU/ml) of haemolytic bacteria (A+B) and non-haemolytical bacteria (C+D). Symbols as for Figure 1. Each data point is plotted and curves outline the mean for each group. Vertical dotted line indicates the last day of pplgG administration. Pairwise comparisons between groups on each day were tested for statistical significance using Mann-Whitney test: a = IgG vs. control (p<0.02).

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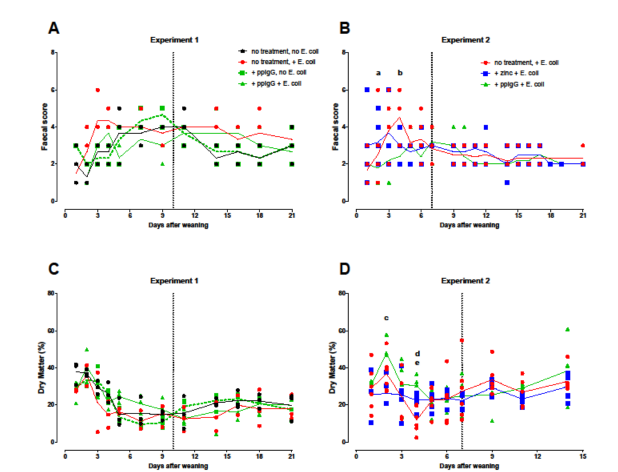
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Figures

Figure 1



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Figure 2

