Vaccination of Dams Increases Antibody Titer and Improves Growth Parameters in Finisher Pigs Subclinically Infected with Porcine Circovirus Type 2

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Porcine circovirus type 2 (PCV2) is the obligate infectious agent in postweaning multisystemic wasting syndrome (PMWS) of pigs. To control PMWS, we vaccinated dams at 4 and 2 weeks before pregnancy and again in the 12th week of gestation with an inactivated PCV2 vaccine (Circovac). Two producer farms run under the control of Swiss Swine Health Organization were selected for the experiment. Previously, in one farm PMWS was diagnosed on pigs after weaning, whereas in the other farm, pigs wasted during the fattening period. For the experiments 113 dams were randomly vaccinated, and 111 dams were sham injected. Vaccination increased serum antibodies in dams 3- to 9-fold, accompanied by serum antibody titer increases in their offspring. In the sixth week of life, progeny from vaccinated dams had about the same IgG antibody titer as progeny of unvaccinated dams at the third day of life. In sera of vaccinated dams only low concentrations of PCV2 DNA were detected, and no progeny developed PMWS. Interestingly, at day 56 four progeny of unvaccinated dams tested positive for anti-PCV2 IgM antibodies, indicating a primary infection with PCV2. Of economic importance is the observation that progeny of vaccinated dams had a significantly higher daily weight gain in the fattening period (farm X, +51 g/day; farm Y, +30 g/day) and thus a shortened fattening period of about 6 days compared to progeny of controls. To our knowledge this is the first demonstration of subclinical circovirus infection and its effects on growth performance of fattening pigs by vaccination of dams.

Postweaning multisystemic wasting syndrome (PMWS) in pigs was first described in Canada (18) and has since been recognized as one of the economically most important swine diseases worldwide (2, 9, 19, 21, 24, 44). PMWS emerged as an epizootic disease in Switzerland in 2003 to 2004 even though cofactors described as important for PMWS development, including porcine reproductive and respiratory syndrome (PRRS), enzootic pneumonia (EP), actinobazillosis, and progressive atrophic rhinitis (pRA), were not present (54).

PMWS is an acute or chronic disease affecting animals at the age of 5 to 16 weeks (1, 11) or exceptionally until 30 weeks of age (37). Typical signs are wasting, profuse diarrhea, and dyspnea, and pigs may have gastric ulcers, enlarged lymph nodes, anemia, icterus, hemorrhages, vasculitis, or edema in various organs (1, 18, 39, 42, 43).

Various porcine circovirus type 2 (PCV2) genotype group members have the potential to be involved in the PMWS etiology (9, 19, 21, 24, 39, 44). Nevertheless, PCV2 can be detected in healthy pigs or isolated from various cells and organs, including peripheral blood, mononuclear cells, dendritic cells, and lymphocytes, and viral antigen is often found in defined lymphatic areas in lymph nodes, tonsils, spleen, and thymus (3, 4) or is scattered in their supporting reticular cells, associated with irregular tissue architecture and in macrophages (39, 49). In other cases, PCV2 was diagnosed in lung, liver, kidney, and the gastrointestinal tract and, in rare cases, in apoptotic vascular endothelial cells of the brain (55).

As PCV2 can replicate in multiple cells of various organs to measurable titers in clinically healthy or diseased animals, the virus may be present in serum or all other body fluids (1, 43) including semen (30, 41). Infection of naïve animals may occur by direct contact with infected animals and their secretions; airborne dissemination must be considered due to high viral loads in large farms (26). In addition, natural vertical transmission was diagnosed in field cases (20, 53) and could be induced experimentally (33, 40). Experimentally infected dams delivered dead and stillborn piglets. PCV2 infection in fetuses was verified and was associated with myocarditis, fibrosis, and degeneration of the myocardium as well as depletion of lymphocytes (32, 38). Recent evidence further suggests that intrauterine infection may have been underestimated at least in some herds (45).

In a retrospective epidemiological study, PCV2 could be traced back to 1979 in Switzerland (54). Nevertheless, the first PMWS case was not confirmed until 2001 (5). However, the epizooty started in late 2003 in areas with large swine populations (52).

PCV2 has been endemic worldwide since the mid-1990s and can be isolated from PMWS-diseased and clinically healthy animals. PCV2-specific antibodies are detected in almost all pigs (1, 16, 29, 36, 48, 51). Another issue is the observation that the profiles of PCV2 serum antibody titers of pigs from
PMWS-affected and unaffected herds are almost identical (17, 23). Thus, the presence of PCV2-specific IgG antibodies is of limited diagnostic or prognostic value and should be considered for diagnostics only in conjunction with disease pattern and PCV2 viral load (46).

Until recently, the main effort to reduce PMWS focused on optimizing herd management in general (27, 28) and intensified monitoring of health status such as the program run by the Swiss Swine Health Organization (www.suisag.ch/SGD/Richtlinien). Despite the absence of porcine reproductive and respiratory syndrome, enzootic pneumonia, actinobazilliosis, and progressive atrophic rhinitis in Switzerland, PMWS occurred in such herds (54). Hence, other measures had to be taken to control the disease. Vaccination against PCV2 was thus considered.

Two types of vaccines against PCV2 were introduced in Europe. One is used to vaccinate pregnant sows to increase colostral antibody concentration while the other is used to vaccinate piglets. Several field studies have demonstrated that vaccination of nursing piglets is effective in reducing losses caused by PMWS and that maternal antibodies present at the time of vaccination did not interfere with active antibody production (13, 22, 35).

In the present study, dams from two different farms under the control of the Swiss Swine Health Organization and with a PMWS history were immunized. The ubiquitous presence of PCV2 may facilitate intrauterine or perinatal infection (8). Vaccination of dams against PCV2 may decrease overall viral load perinatally, and increasing colostral antibodies may protect offspring within the first days of life. To test the effectiveness of vaccination on reproductive parameters, antibody production of the dams and antibody transfer to piglets, mortality rate, growth performance of offspring, and age of slaughter were analyzed.

MATERIALS AND METHODS

History of the herds. This study took 14 months. We used Cirovac (Merial SA, Lyon, France) in two different farms with a history of recurrent PMWS. In both herds PMWS had been diagnosed before birth of the first litter. PMWS diagnosis was according to Sorden (46) and described by Wiederkircher et al. (54).

Herd X was a breeding herd with 90 Swiss Large White sows. Gilts were raised for replacements within the farm. Breeding dams and weaned pigs were kept in different barns on the same farm. Weaned pigs were sold to a regional finisher at the age of approximately 10 weeks at a body weight of 22 to 27 kg. Only cross-fostering occurred; however, these dams were categorized as either young (≤3 litters; n = 65) or experienced (>3 litters; n = 159), as indicated in the data sets. Thus, the farm was in a subclinical-infection phase. During the study no changes were undertaken, too. When PMWS was suspected, tissues were examined immunohistochemically (IHC) and with PCR for PCV2 infection (31, 47, 54). Additionally, on farm Y, 2 to 5 ml of blood was collected from each of 101 individually tagged, randomly selected piglets from 17 litters at the age of 3, 10, 31, 42, 56, and 63 days postpartum (pp). Our study was carried out according to Swiss Animal Welfare guidelines (study number 06/07).

Serological examinations. A competitive enzyme-linked immunosorbent assay ([ELISA] SerELISA PCV2 AB Mono Blocking Systems; Symbiotics Corporation Europe SAS, Lyon, France) was used for antibody (IgG) detection (14). The completion of the test, data analysis, and transformation of the data into ELISA units (EU) were done according to the manufacturer's instructions and a published reference (14). We supplemented the assay with two additional controls to check quality to serum dilutions suggested by the manufacturer. First, we used additional positive- and negative-control sera to check plate antigen coating homogeneity. Second, we normalized S values (linear s/n ratio [14]) among individual plates with the aid of a known serum. Immunoglobulin M (IgM) was measured using Ingezim Circovirus IgG/IgM (Ingenasa, Madrid), a capture immunoenzymatic assay specific for IgM antibody detection to PCV2.

Production variables of sows. Parity number, litter weight, number of live- and dead-born piglets, number of mummified pigs, number of piglets weighing below 1 kg, number of piglets lost during the nursing period, and cause of death were recorded for each dam.

Production variables of progeny. Cross-fostering occurred; however, these piglets were excluded. The number of finishing days was calculated from the dates of birth and slaughter (ADWG 2) is calculated as live slaughter weight (kg) divided by age in days. The average daily weight gain in the fattening period (ADWG 3) is calculated as live slaughter weight (kg) minus the weight at the beginning of the finishing period (kg) divided by the number of finishing days. The carcass weight considered to represent 78% of the live weight at slaughter and was used to calculate the latter. The number of finishing days was calculated from the dates of weaning and slaughter.

Pathological examinations. All mummified and stillborn pigs and dead nursing piglets as well as the dead weaning and fattening pigs were examined at the Institute of Veterinary Pathology, Vetsuisse Faculty, University of Zurich. If deemed necessary, histological, bacteriological, and virological examinations were undertaken, too. When PMWS was suspected, tissues were examined immunohistochemically (IHC) and with PCR for PCV2 infection (31, 47, 54).

Statistics. Statistical calculations were carried out using StatView, version 5.1 (SAS Corporation). Analysis of variance (ANOVA) for repeated measures and unpaired and paired t tests were considered statistically significant at a P value of ≤0.05.

RESULTS

Production parameter of dams. This study was conducted within a period of 14 months. Dam production parameters, including gestation rate, numbers of mummified, aborted, or weak-born and live-born piglets, and litter size, did not change during the 14 months and did not differ between vaccinated and unvaccinated dams. Also, there were no piglet loss differences observed between the vaccinated and nonvaccinated dam progeny during suckling or weaning time.

Postmortem examination. Of a total of 2,720 piglets born, 379 (13.9%) died and underwent a postmortem examination. Of these, 176 were from herd X, and 203 were from herd Y. This number includes aborted and stillborn piglets and perinatal losses. The majority of these piglets were crushed by their
mothers during the first week of life. Sixty of the dead piglets were examined immunohistochemically for the presence of PCV2 antigen. All were negative. At the weaning stage, 9 of 1,169 weaned pigs died in herd X (0.8%), and 15 of 1,172 weaned pigs died in herd Y (1.3%). In none of these pigs was PMWS diagnosed.

In the fatten period PMWS was diagnosed in 2 of the 112 pigs born to two nonvaccinated sows of herd X (1.8%). Interestingly, no case of PMWS was diagnosed in 311 fatten pigs from herd Y.

**Anti-PCV2 IgG antibody titer of dams and PCV2 infection.**

At the start of the experiment (time B0), 99% of all dams had antibodies against PCV2. In herd X, the mean titer at B0 of all sows was significantly ($P = 0.006$) higher than that of herd Y (Table 1). At time points B1 and B2, vaccinated experienced as well as young dams of both herds (Table 1) had significantly higher titers against PCV2 than unvaccinated dams ($P \leq 0.05$).

Upon vaccination, antibody titers increased up to 9-fold.

We noted a fluctuation in titers over time in the unvaccinated dams that may have been due to subclinical infections. The dams of herd Y had a lower titer at time point B0 than herd X (Table 1). The unvaccinated controls showed a mild but significant increase in titer (herd X, $P = 0.03$; herd Y, $P = 0.04$) between time points B0 and B1 and a decrease between B1 and B2.

A closer analysis of the titers of young and experienced dams revealed some interesting observations (Table 1). Young dams had a 3.5-fold higher baseline titer (B0) than experienced dams ($P < 0.0001$). At B1, vaccinated young dams had the highest titer (6,820 EU) measured during the entire study. Only low concentrations of PCV2 DNA (<5 x 10^7 copies/ml) were detected by PCR in some sera (data not shown).

**Offspring anti-PCV2 IgG and IgM antibody titers.** For experiments determining the anti-PCV2 IgG and IgM antibody titers of offspring, sera from 101 randomly selected offspring of vaccinated and unvaccinated dams were collected in herd Y. At all times, the piglets from vaccinated dams had significantly higher PCV2 antibody titers than piglets from unvaccinated dams ($P < 0.001$) (Table 2 and Figure 1). From day 3 postpartum (pp) to day 63 pp, the antibody titer in the vaccinated group decreased gradually from 5,173 EU to 511 EU. In the piglets from the unvaccinated dams, the antibody level decreased from 2,323 EU at day 3 pp to 86 EU at day 63 pp. The decrease was gradual until day 63. At no time point was a seroconversion noted, as previously described (12). At this stage, this is paralleled by the absence of any case of PMWS.

To detect potential subclinical infections, sera of all piglets from unvaccinated and vaccinated dams collected at days 10, 31, 42, 56, and 63 were examined for IgM antibody against PCV2. Four piglets from unvaccinated sows in herd Y were positive at day 56, indicative of subclinical infections.

**Offspring production parameters.** During the fatten period, a total of 423 randomly selected pigs from 218 litters were monitored until slaughter (112 of herd X and 311 of herd Y). Offspring of vaccinated dams from both herds X and Y had significantly greater average daily weight gains from birth to slaughter (ADWGs) than the controls of unvaccinated dams (Table 3). The difference in ADWG values between progeny from vaccinated and unvaccinated dams was 33 g/day in herd X and 20 g/day in herd Y. The difference in the average daily weight gains in the fatten period (ADWG’s) between offspring of vaccinated and unvaccinated dams was 51 g/day in farm X and 30 g/day in farm Y. The age at slaughter of pigs from vaccinated dams was reduced by 6.7 days in herd X ($P = 0.03$) and by 5.5 days in herd Y ($P = 0.02$). Therefore, the increased antibody titers in dams were transmitted by colostrum and were associated with a possible impact on progeny health status, as reflected by increased weight gain in the fatten period.

**DISCUSSION**

Neither farm showed any reproductive disorders due to PCV2. The perinatal loss of some 14% was not affected by vaccination, and this value is in the range of small to mid-sized pig farms surveyed by the Swiss Swine Health Organization (50). Therefore, the existing natural immune responses of the dams against PCV2 enhanced by vaccination appeared sufficient to prevent fatal intrauterine infection. This is not surprising as PCV2-caused reproductive disorders are infrequent (16). Furthermore, all collected sera from adult animals

### Table 1. Antibody titers of dams in farm X and Y and of young and experienced dams

| Time   | All herd X dams | All herd Y dams | Young dams | Experienced dams |
|--------|-----------------|-----------------|------------|------------------|
|        | Vaccinated      | Unvaccinated    | Vaccinated | Unvaccinated     |
|        | $(n = 67)$      | $(n = 68)$      | $(n = 46)$ | $(n = 43)$       |
| B0     | 1.613 ± 306     | 1,893 ± 369     | 53) Unvaccinated | $(n = 53)$ |
|        | 3.055 ± 410     | 2,140 ± 339     | $<0.05$ | $<0.05$ |
|        | 5,156 ± 358     | 1,670 ± 344     | $<0.05$ | $<0.05$ |
| Values are mean antibody titers and standard deviations.

### Table 2. Antibody titers of piglets from vaccinated and unvaccinated dams of herd Y

| No. of days postpartum | Antibody titer (EU)$^a$ | $P$ value |
|------------------------|------------------------|----------|
|                        | Vaccinated $(n = 53)$  | Unvaccinated $(n = 48)$ |
| 3                      | 5,173 ± 322            | 2,323 ± 308 | $<0.05$ |
| 10                     | 4,319 ± 248            | 1,266 ± 203 | $<0.05$ |
| 31                     | 3,062 ± 359            | 1,037 ± 206 | $<0.05$ |
| 42                     | 1,943 ± 296            | 279 ± 69   | $<0.05$ |
| 63                     | 511 ± 81               | 86 ± 31    | $<0.05$ |

$^a$ Values are mean antibody titers and standard deviations.
showed low concentrations of PCV2 DNA (<5 × 10^5 copies/ml). Also, with the first litter important for this study, we did not find any indication of PCV2-associated diseases in 379 analyzed dead pigs. In contrast, the historical loss of 5 to 10% of pigs after weaning and in the early fattening period compared to data points of interest had improved 6- to 12-fold. Only 2 of the 423 evaluated fattening pigs died of PMWS during the experiment in the early finishing period. Interestingly, the two pigs in question were born to unvaccinated dams. Therefore, we suggest that the study be conducted in a postepizootic period of the PCV2 disease cycle.

As expected due to previous epidemiological and serological examinations, all dams had antibodies from natural exposure to PCV2 before vaccination (23, 26). In contrast to other investigators (34, 35), we opted to vaccinate twice before and once during pregnancy to compensate for a potential loss of serum antibodies into the colostrum (7) and to improve reproduction parameters. Antibody titers were boosted 3- to 9-fold after two vaccinations, and these values either leveled off or decreased in spite of the third immunization. Decreased antibody titers were also observed in unvaccinated dams. This is an independent indication that the serum antibody decline may be attributed to colostrum antibody supplementation that could not be overcome by additional vaccination.

Right after birth, PCV2-specific antibody titers in sera from piglets of vaccinated dams were very similar to those of their mothers, as previously described (7). The antibody titers in these piglets decreased gradually about 10-fold within 60 days and was 2.5 to 6 times higher than that of the controls at the same point of time. Antibody titers against PCV2 in neonates of unvaccinated dams reflected those of their mothers in midgestation (7), but the decrease in the mean titers in these animals until day 63 was 30-fold. The steeper proportional antibody loss in the control piglets might be because their anti-PCV2 antibodies are less mature than the anti-PCV2 antibodies elicited by vaccination.

It is very difficult to induce a protective immune response in mammals right after birth. Therefore, piglets are vaccinated against PCV2 at the weaning stage. In the first weeks of life, piglets are protected by maternal antibodies that may vary depending on the immune status of the dam. Thus, piglets will be exposed to the PCV2 viral pressure present in the herd, with potentially devastating effects. The vaccination of dams appeared to decrease the infective pressure generally and provided increased antibodies against PCV2 in colostrum to protect piglets within the first days of life. At this age colostral antibody is the only specific immune mediator available to the piglets until they can generate their own. We calculated by linear regression of anti-PCV2 IgG antibody concentrations that progeny antibody titers from vaccinated dams at day 49 pp were about the same as the titers of the controls at day 14 pp. Interestingly enough, we found that 4 of 48 pigs from the unvaccinated group contained IgM antibodies against PCV2 at the age of 56 days pp. Based on the IgM antibodies observed, we concluded either that these four piglets were newly PCV2 infected, probably around day 49 pp, or that the antibodies of this isotype were not detected in previous testing (data not shown). Nevertheless, we could not detect PCV2 DNA in blood of these piglets (data not shown). The protective anti-

![FIG. 1. Piglet anti-PCV2 IgG antibody titers. Blood (2 to 5 ml) was collected from offspring of vaccinated (filled black bars) or unvaccinated (filled gray bars) dams at the age of 3, 10, 31, 42, and 63 days postpartum (pp). Anti-PCV2 IgG concentrations were determined by ELISA. At all times, the piglets from vaccinated dams had significantly higher anti-PCV2 antibody titers than piglets from unvaccinated dams (*, P < 0.001).

### TABLE 3. Production parameters of offspring from vaccinated and unvaccinated dams at farm X and farm Y

| Parametera | Farm X progeny (n = 112) | Farm Y progeny (n = 311) |
|------------|--------------------------|--------------------------|
|            | Vaccinated (n = 57) | Unvaccinated (n = 55) | P value | Vaccinated (n = 182) | Unvaccinated (n = 119) | P value |
| ADWG1 (g/day) | 626 ± 0.01 | 593 ± 0.01 | <0.05 | 585 ± 0.01 | 565 ± 0.01 | <0.05 |
| ADWG2 (g/day) | 785 ± 0.02 | 734 ± 0.02 | <0.05 | 726 ± 0.01 | 696 ± 0.01 | <0.05 |
| Age at slaughter (days) | 170 ± 1.9 | 176 ± 2.2 | <0.05 | 183 ± 1.8 | 189 ± 1.4 | <0.05 |

a ADWG1, average daily weight gain, calculated as live slaughter weight (kg) divided by age in days; ADWG2, average daily weight gain in the fattening period, calculated as live slaughter weight (kg) minus weight at the beginning of the finishing period (kg) divided by the number of finishing days.
body titers at day 14 pp compare to values from the literature as PMWS is not found before 3 weeks of age.

The general health status improvement of pigs from vaccinated dams was further observed in the significantly high average daily weight gain and decreased time to slaughter compared to offspring from unvaccinated dams. It is tempting to speculate that the improved immune status of vaccinated dams and the increased collostral antibodies taken up by their offspring decreased the overall infectious pressure and enhanced overall viability of the progeny. Yet a direct comparison of results in an industrial setting is difficult. Vaccination of dams as done here, rather than on 2- to 3-week-old piglets (10, 12, 22), might prove advantageous. The perinatal virus load of gilts or piglets may have been underestimated at least in some herds (15, 45). PCV2 can infect a variety of cells and preferentially replicates in dividing cells (49). Dividing cells are very important to the process of reaching adult stage cell numbers; they are crucial for the development of the architecture of organs of the immune system, the gastrointestinal tract, and the central nervous system. Failure or delayed maturation of the immune system may favor secondary infections or growth retardation due to inadequate food intake or inadequate hormonal regulation of body growth (6). The weight gain of young animals may thus be considered a mirror reflecting the combined effect of a virus on rapidly dividing cells necessary for individual development.

This is the first evidence of effective vaccination in a PCV2 subclinical infection as determined by daily weight increase of pigs during the fattening period. Dam vaccination was associated with offspring health status improvement at the weaning stage, which became obvious in the fattening period in PMWS postepizootic farms. Offspring of vaccinated dams outperformed pigs from unvaccinated dams in terms of weight gain and decreased maturity time for slaughter in the fattening period. Even in the absence of overt PCV2-associated diseases, small amounts of PCV2 might interfere with the maturation of individuals, possibly by interfering with rapidly diving cells. We associated these economic benefits with low PCV2 viral pressure due to vaccination of dams and their transfer of collostral antibodies to progeny in particular.

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