Oral antigen exposure in newborn piglets circumvents induction of oral tolerance in response to intraperitoneal vaccination in later life

J Alex Pasternak1, Siew Hon Ng1, Rachelle M Buchanan1, Sonja Mertins2, George K Mutwiri1, Volker Gerdts1 and Heather L Wilson1*

Abstract

Background: We previously determined that newborn piglets orally gavaged with Ovalbumin (OVA) responded to systemic OVA re-exposure with tolerance; if adjuvants were included in oral vaccine, piglets responded with antibody-mediated immunity (Vet Immunol Immunopathol 161(3–4):211–21, 2014). Here, we will investigate whether newborn piglets gavaged with a vaccine comprised of OVA plus unmethylated CpG oligodeoxynucleotides (CpG; soluble component; OVA/CpG) combined with OVA plus CpG encapsulated within polyphosphazene microparticles (MP; particulate component) responded with systemic and mucosal immunity. To monitor the response to systemic antigen re-exposure, piglets were i.p.-immunized with OVA plus Incomplete Freund’s Adjuvant (IFA) one month later.

Results: Newborn piglets (n = 5/group) were gavaged with a combined soluble and particulate vaccine consisting of OVA (0.5–0.05 mg) plus 50 μg CpG and 0.5 mg OVA plus 50 μg CpG encapsulated within a polyphosphazene MP (0.5 mg) referred to as OVA/CpG + MP. Control piglets were gavaged with saline alone. Piglets were i.p. immunized with 10 mg OVA (or saline) in IFA at four weeks of age and then euthanized at eight weeks of age. We observed significantly higher titres of serum anti-OVA immunoglobulin (Ig) IgM, IgA, IgG, IgG1, IgG2 and IgG in piglets immunized with 0.05 mg OVA/CpG + MP relative to saline control animals. Thus, a single oral exposure at birth to a combined soluble and particulate OVA vaccine including adjuvants can circumvent induction of oral tolerance which impacts response to i.p. vaccination in later life. Further, piglets gavaged with 0.05 mg OVA/CpG + MP generated significant anti-OVA IgG and IgG1 titres in lung compared to saline control piglets but results were comparable to titres measured in parenteral control piglets. Peripheral blood mononuclear cells (PBMCs) ex vivo-stimulated with OVA showed markedly decreased production of IL-10 cytokine after 72 hours relative to animal-matched cells incubated with media alone. No production of IFN-γ was observed from any groups.

Conclusion: Newborn piglets gavaged with low dose soluble and particulate OVA plus CpG ODN and polyphosphazene adjuvants produced antigen-specific antibodies in serum and lung after systemic re-exposure in later life. These data indicate circumvention of oral tolerance but not induction of oral immunity.

Keywords: Piglets, Neonate, Ovalbumin, Oral, Tolerance

* Correspondence: heather.wilson@usask.ca
1Vaccine and Infectious Disease Organization (VIDO), University of Saskatchewan, 120 Veterinary Road, Saskatoon, SK S7N 5E3, Canada
Full list of author information is available at the end of the article

© 2015 Pasternak et al; licensee BioMed Central. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly credited. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background
If exposure to an antigen by the oral route fails to promote an oral immune response, any subsequent re-exposure (even by a systemic route) results in suppression of immunity; this process is known as oral tolerance. Oral tolerance is a major suppressive immunological process designed to prevent local and peripheral over-reaction to innocuous antigens [1,2]. Commensal bacteria are critically required for proper gut-associated lymphoid tissue (GALT) development and induction of oral tolerance [3,4]. Mucosal dendritic cells (DCs) play an active role in inducing oral tolerance through mechanisms which require retinoic acid, vitamin D, interleukin (IL)-10, Transforming growth factor (TGF)-β, and indoleamine-2,3-dioxygenase [5-9]. In the mesenteric lymph nodes (MLNs), T regulatory (Treg) cells undergo differentiation and home back to the inductor site to induce and/or maintain antigen-specific oral tolerance [8]. Several physical barriers prevent antigen/pathogen contact with GALT and subsequent penetration of the gut wall making targeted induction of oral immunity a significant challenge [10,11]. The gut of the newborn piglet is uniquely designed to be semi-permeable or ‘leaky’ for a limited time to allow colostrum-derived cells, antibodies, and other macromolecules such as albumin, cytokines, antimicrobial peptides and many other bioactive products to be passively transferred to the piglets. These maternally-derived cells and macromolecules traverse the gut wall then enter into the vasculature where they play a variety of roles including passive immunity against disease [12-15]. ‘Gut closure’ occurs within a few days after birth in ruminants [16] and pigs [17], but it does not occur until after weaning (two weeks of age) in rats and mice [18-20]. In humans, a considerable amount of ‘gut-closure’ occurs both before birth and within a few days after birth but it may in fact take up to two years to reach the same level of impermeability that is observed in the adult gut [21,22]. Once across the gut wall, antigens may be able to interact with DCs within the sub-epithelial dome which can then present antigens to T cells within isolated lymphoid follicles and/or Peyer’s Patches to promote induction of oral immunity rather than being taken up by tolerogenic mucosal DCs which promote oral tolerance [23,24]. Despite the overwhelming propensity to respond to an oral antigen with tolerance, oral vaccines are highly sought because of their ease of administration. They are needle-free and therefore present reduced risk of transmitting infections and less need for qualified personnel to administer the vaccine. Moreover, an estimated 90% of all pathogens invade through mucosal surfaces, therefore mucosal immunity (induced by mucosal vaccines) offer the potential to control pathogens at their point of entry.

Previous work from our lab showed that rat pups and lambs orally vaccinated starting the day after birth with multiple doses of soluble ovalbumin (OVA; without adjuvants) responded with immunity to subsequent intraperitoneal (i.p.) immunization [25,26]. In contrast, we showed that newborn piglets orally immunized within six hours after birth with single bolus of soluble OVA then boosted through the i.p. route one month later showed significantly lower anti-OVA immunoglobulin (lg) A titres and a strong trend towards lower anti-OVA IgM, IgG1, IgG2 and IgG titres relative to the i.p. control group indicating induction of oral tolerance [27]. These data showed agreement with Haverson et al [28] who demonstrated that newborn piglets orally vaccinated once with OVA induced classical oral tolerance following a systemic challenge by showing reduced specific systemic IgG responses. When we included unmethylated oligonucleotides containing CG oligodeoxynucleotides (CpG ODNs) and soluble polyphosphazene in the oral vaccine administered at birth, the response to i.p. immunization one month later was induction of immunity (i.e. increased serum anti-OVA IgA, IgM, IgG1, IgG2 and IgG titres relative to piglets immunized with OVA alone) [27]. Clearly the components of the oral vaccine administered at birth impacted the response to the booster immunization one month later.

Factors contributing to induction of oral tolerance include: the host’s immunological maturity at time of exposure, the timing and the frequency of exposure, and the nature of the antigen [29-32]. We established that a single bolus of soluble OVA with CpG ODN and polyphosphazene adjuvants administered the day after birth induced oral immunity in piglets [27]. Our next step will be to establish whether inclusion of the antigen in a particulate form promotes oral immunity and whether the response could be observed at distal mucosal sites. To test this hypothesis, conventionally reared neonatal piglets were gavaged within six hours after birth with 0.5 mg or 0.05 mg OVA plus CpG ODN in a soluble form as well as OVA plus CpG ODN encapsulated within a polyphosphazene microparticle (MP) [33-35]. Systemic and mucosal antibody titres and ex vivo cytokine production are assessed to determine antibody-mediated and cell-mediated immunity

Results
Explanation of vaccination procedure
Piglets were gavaged with a two-part vaccine consisting of soluble OVA (0.5 mg or 0.05 mg) with 50 μg soluble CpG 2395 (which together make up the soluble components of the vaccine; OVA/CpG) as well as a MP encapsulating 0.5 mg OVA + 50 μg CpG 2395 (which comprised the particulate part of the vaccine). These vaccines will be referred to as 0.5 mg OVA/CpG + MP
and 0.05 mg OVA/CpG + MP. The experimental timeline is detailed in Figure 1. Piglets were gavaged at less than six hours of age when their gut would be semi-permeable with the idea that both the soluble OVA/CpG and the OVA and CpG ODN within the MP would cross the leaky gut wall. Polyphosphazene-based MPs are water-soluble and would then dissolve over time to release the encapsulated OVA and CpG ODN thus acting like a prime-boost [36,37]. Piglets were bled three days after birth, day seven after birth, and weekly thereafter (Figure 1, grey arrows specify bleed times). Piglets were i.p. immunized with 10 mg OVA (or saline) plus Incomplete Freund’s Adjuvant (IFA) at 28 days of age and all piglets were euthanized at 49 days of age. The i.p. control group received a saline gavage but piglets were boosted with the i.p. vaccination (OVA + IFA) to act as our primary i.p. vaccine control group.

Newborn piglets vaccinated by oral gavage responded with significant serum anti-OVA IgM, IgA, and IgG1, IgG2 and IgG production after re-exposure by the i.p. route

The definition for oral tolerance is that oral exposure to antigen which is subsequently encountered via a systemic route triggers reduced immune responses (such as antibody production) relative to animals exposed to antigen systemically without prior oral exposure [38]. Previous work in our laboratory showed that animals orally immunized six hours after birth with 5 mg or 0.05 mg OVA responded to i.p. immunization at one month of age with oral tolerance, not oral immunity [27]. Therefore these groups (OVA alone without adjuvants) were not repeated here. In the current trial, when we assessed the serum antibody titres in the piglets prior to weaning (less than day 21) for all groups, we observed negligible anti-OVA antibodies of any isotype indicating that the sows did not pass any interfering OVA-specific passive immunity to the piglets (data not shown). Even one week after weaning (Figure 2, day 28), all isotypes of piglet serum anti-OVA antibodies titres were negligible suggesting that oral gavage at birth with 0.5 mg or 0.05 mg OVA/CpG + MP alone did not promote antibody-mediated immunity. On day 42, which was two weeks after the i.p. booster immunization, we observed a significant increase in serum anti-OVA IgM (Figure 2A; p < 0.05), IgA (Figure 2B; p < 0.05), IgG1 (Figure 2D; p < 0.05), and IgG2 (Figure 2E; p < 0.05) titres in the i.p. control group relative to the saline control group. The group gavaged with 0.05 mg OVA/CpG + MP showed significant induction of anti-OVA IgM (Figure 2A; p < 0.05), IgA (Figure 2B; p < 0.01), IgG (Figure 2C; p < 0.05), IgG1 (Figure 2D; p < 0.01), and IgG2 (Figure 2E; p < 0.01) relative to the saline control group suggesting that prior oral exposure to low dose OVA circumvented induction of oral tolerance. The animals gavaged with ten-fold higher dose of soluble OVA (0.5 mg OVA/CpG + MP) showed a strong trend towards increased anti-OVA antibody production relative to the saline control group but the data were not statistically significant.

On day 49, serum titres from animals within the i.p. control group showed a significant increase in anti-OVA IgG1 antibodies (Figure 2D; p < 0.05) relative to the saline control group. In contrast, the group gavaged with 0.05 mg OVA/CpG + MP showed significantly more anti-OVA IgM (Figure 2A; p < 0.05), IgA (Figure 2B; p < 0.05), IgG1 (Figure 2D; p < 0.01), and IgG2 (Figure 2E; p < 0.01) titres than what was observed in the saline control group. The group gavaged at birth with 0.5 mg OVA/CpG + MP showed significantly more anti-OVA

![Figure 1 Description and timeline of immunization protocol. Piglets (n=5/group) were gavaged with ovalbumin (OVA) (0.05 mg or 0.5 mg) + 50 μg unmethylated oligonucleotides containing CG oligodeoxynucleotides (CpG 2395) as a soluble vaccine as well as with 0.5 mg OVA + 50 μg CpG 2395 within a 0.5 mg PCEP polyphosphazene microparticles (MP). Gavages took place within six hours of birth. The negative control (saline) group and the i.p. control group were gavaged with saline. With the exception of the saline control group, the remaining groups were i.p. immunized at four weeks of age with 10 mg OVA in Incomplete Freund’s Adjuvant (IFA). At eight weeks of age, piglets were euthanized and lung lavages were harvested. Blood was obtained on day three, day seven and then weekly (grey arrows). At time of death, blood was drawn for PBMC isolation.](https://example.com/figure1.png)
Figure 2 (See legend on next page.)
IgG1 (Figure 2D; p < 0.05), and IgG2 (Figure 2E; p < 0.05) antibodies compared to the saline control group, but the other isotypes did not. Thus, unlike our previous data which showed that piglets orally vaccinated with soluble OVA alone induced oral tolerance [27], data from the current trial shows that piglets orally vaccinated with 0.5 mg or 0.05 mg OVA/CpG + MP showed significant induction of anti-OVA antibodies in serum indicating circumvention of oral tolerance.

Newborn piglets vaccinated orally with low dose OVA/CpG + MP responded with significant anti-OVA IgG1 and IgG titres in lung lavage after re-exposure by the i.p. route

According to the Common Mucosal Immune System theory, antigen-sensitized precursor B and T lymphocytes generated at one mucosal site (i.e. such as the gut) can be detected at anatomically remote and functionally distinct compartments (such as the respiratory mucosa) [39-45]. After 49 days, the pigs were euthanized and bronchoalveolar lavage was collected. Piglets gavaged with saline but injected with OVA and IFA by the i.p. route alone (i.e. the i.p. control group) failed to trigger significant anti-OVA IgM (Figure 3A), IgA (Figure 3B), IgG (Figure 3C), IgG1 (Figure 3D), or IgG (Figure 3E) titres relative to the saline control group. Newborn piglets gavaged with 0.05 mg OVA/CpG + MP had very low titres of anti-OVA IgM (Figure 3A), IgA (Figure 3B), and IgG2 (Figure 3E) but the titres for anti-OVA IgG (Figure 3C, p < 0.05) and IgG1 (Figure 3D; p < 0.05) were statistically higher than the saline control group. Collectively, these results suggest that oral exposure to OVA/CpG + MP triggered low level mucosal immunity at a distal site.

Neonatal piglets gavaged with OVA/CpG + MP did not respond with induction of OVA-specific cell-mediated immunity

Finally, we sought to determine whether neonatal piglets orally gavaged with 0.5 mg or 0.05 mg OVA/CpG + MP developed cell-mediated immunity as measured by IFN-γ (Type 1 T-helper cell (Th1)) cytokines and IL-10 (Th2 cytokines). Peripheral blood mononuclear cells (PBMCs) were collected at eight weeks of age and they were restimulated ex vivo with OVA or media for 72 hours before the supernatants were collected and antibody titres were assessed. PBMCs did not show OVA-specific induction of IFN-γ for any group except in the presence of the mitogen Concanavalin A (data not shown). Interestingly, we observed a decrease in production of the anti-inflammatory cytokine IL-10 relative to unstimulated (media control) cells in all groups including the saline control groups. These data indicate that oral gavage of newborn piglets did not promote cell-mediated immunity as measured at 8 weeks of age.

Discussion

Because the vast majority of infectious agents enter the body through mucosal routes, it is reasonable to assume that mucosal immunity which combats the infectious agent prior to colonization and penetration would be much more effective than systemic immunity. But it has proven very challenging to design effective oral subunit vaccine without the use of very strong mucosal adjuvants such as cholera toxin [46,47]. The majority of clinically approved oral vaccines for use in pigs are live attenuated viruses or bacteria (www.vetvac.org/index.php). One may speculate that in order to trigger an immune response instead of tolerance, the pathogen must traverse the gut wall and/or penetrate the epithelial cells lining the gut wall. If this is the case, it is understandable that subunit vaccines that lack strong adjuvants such as cholera toxin should fail to promote immunity [48]. However, such adjuvants cause significant side effects and are therefore not in clinical use. Due to the inherent risk of attenuated pathogen vaccines reverting to virulence, live attenuated vaccines against pathogens such as Porcine Reproductive and Respiratory Virus are not recommended for use in seronegative herds and are therefore a reactive vaccine instead of a proactive vaccine [49-51].

Previous work in our lab showed that oral administration of soluble OVA in rat pups or lambs starting immediately after birth was sufficient to promote oral immunity or at minimum prevented induction of oral tolerance [25,26]. In contrast, our research with piglets showed that soluble OVA administered immediately after birth triggered induction of oral tolerance [27]. However, if CpG ODN and polyphosphazene adjuvants were included in the soluble oral vaccine, there was instead evidence of induction of serum antibody-mediated immunity in response to systemic re-exposure in later life [27]. In the current study, we investigated whether oral immunization with a mixture of soluble and particulate OVA plus CpG ODN and polyphosphazene adjuvants could promote oral
Figure 3 (See legend on next page.)
immunity. We determined that systemic anti-OVA antibody-mediated immune responses were induced in newborn piglets gavaged once on the day of birth with 0.5 mg or 0.05 mg OVA/CpG + MP and subsequently boosted with OVA and IFA by the i.p. route one month later. If oral tolerance to OVA had been induced instead, re-exposure to the antigen by i.p. injection should have resulted in a reduction of serum anti-OVA antibody titres as observed in [27]. Anti-OVA IgG and IgG1 titres in lung lavages were significantly induced in piglets gavaged at birth with 0.05 mg OVA/CpG + MP indicating induction of mucosal antibody-mediated immunity at a distal mucosal site. IgG has not traditionally been recognized as a major mucosal immunoglobulin, however there is growing evidence that oral vaccines can elevate local and systemic IgG titres [52,53] and that IgG antibodies may play a role in passive transfer of luminal antigens across the gut wall using FcRN [54]. Therefore, our data shows that vaccination of newborn piglets with a joint soluble and particulate subunit vaccine triggered systemic antibody-mediated immunity which contrasts with what is reported in older piglets orally vaccinated with soluble antigens or newborn piglets gavaged with OVA alone [27,52].

Results from our lab showed that oral vaccination of newborn lambs with OVA produced minimal cell-mediated immunity as established by IFN-γ expression and lymphocyte proliferation in ex vivo stimulated spleenocytes [26]. Similarly, ex vivo-stimulated mLN cells from rat pups gavaged with OVA after birth failed to produce significantly higher IFN-γ titres relative to cells from pups gavaged after birth with saline [25]. In the current study, we gavaged piglets with soluble and particulate OVA plus polyphosphazene and CpG ODN, the latter of which is known to promote IFN-γ production [55]. Despite inclusion of CpG ODN in the oral piglet vaccine, IFN-γ production was negligible suggesting that oral immunization of newborns may not induce significant cell-mediated immunity.

Further studies must be undertaken to clarify the precise dose, vaccine formulation and timing of exposure required for induction of cellular immunity as well as including direct measurements of mucosal immunity. Experiments are underway to elucidate the kinetics of gut permeability and the impact this has on the mechanisms of antigen uptake and where antigen presentation to lymphocytes occurs (i.e. in the Peyer’s patches, isolated lymphoid follicles, or mesenteric lymph nodes). Should early life oral vaccination consistently circumvent induction of oral tolerance and/or promote oral immunity, it will have important implications for protecting against infectious diseases in the very young and it may reduce the number of carriers of disease-producing organisms within a herd.

Conclusions
In the present study, we determined that low dose OVA/CpG + MP circumvented induction of oral tolerance with comparable serum anti-OVA antibodies relative to the animals gavaged with 0.05 mg OVA/CpG + MP relative to the i.p. control animals. There was a trend towards induction of mucosal immunity but the results were not statistically significant. These results are intriguing and should be studied further to establish whether it may be advisable to proactively orally vaccinate newborn piglets to prevent induction of oral tolerance to ensure that the they can respond appropriately to parental vaccines in later life.

Methods

Immunization procedure
This work was approved by the University of Saskatchewan’s Animal Research Ethics Board, and adhered to the Canadian Council on Animal Care guidelines for humane animal use. Pregnant Landrace-cross sows were housed at VIDO with ad libitum access to standard feed and water. Piglets were randomly assigned to treatment groups. We gavaged piglets with a single bolus of saline or 0.5 mg or 0.05 mg OVA (Sigma-Aldrich Canada Ltd, Oakville, ON, Canada) plus 50 μg CpG 2395 (soluble components; 5′- TCGTCGTTCGCGCACGC GCCG -3′ phosphorothioate oligodeoxy-nucleotide from Merial Limited (Lyon, France)), as well as the particulate component comprised of 0.5 mg OVA + 50 μg CpG 2395 + 500 μg poly [d(sodiumcarboxylatoethylphenoxy)-phosphazene] (PCEP) (synthesized by Idaho National Laboratory (Idaho Falls, ID, USA). (These doses were extrapolated from a successful oral vaccination with OVA in lambs and weight adjusted [26]). Microparticles were formulated as detailed in [34]. A total volume of
10 mL was administered via gavage using soft Nalgene Tubing with a monojet catheter tip (Fisher Scientific Ltd., Ottawa, ON) gently inserted into the back of the throat. Piglets were bleed on day three and day seven and then weekly until day 49 (Grey arrows, Figure 1). All blood samples were collected using Ethylenediaminetetraacetic acid (EDTA) Vacutainers (BD Biosciences-Canada, Mississauga, ON), centrifuged (4547 x g) and serum stored at -20°C until antibody titres were measured. At four weeks age, piglets were i.p.-injected with 10 mg OVA plus IFA (Sigma-Aldrich). To generate the peripheral control group, piglets received saline by oral gavage and they were i.p. immunized with OVA plus IFA at four weeks as indicated above. This route was used because it is considered relevant for stimulating the mucosal tissues [56,57]. On day 49, piglets were euthanized using 2 mL/10 lb body weight with Pentobarbital Sodium Injection (240 mg/mL; Euthanyl, Bimeda-MTC Animal Health Inc., Cambridge, ON). Lung lavages were obtained at the end of the trial (Eight weeks; Figure 1). The lung lavage was kept on ice until centrifugated at 400 x g at 4°C for 10 min, then cells were washed twice with cold PBS, counted, and suspended to a final concentration of 4 x 10^6 cells/mL in 10% complete Roswell Park Memorial Institute (RPMI) medium (Gibco; Life Technologies, Burlington, Ontario), supplemented with 0.2 mM L-glutamine, 0.1 mM HEPES, 0.05 mg/mL gentamicin, 0.02 mM 2-mercaptopetanol (Sigma-Aldrich for all), and 10% heat inactivated horse serum (Gibco; Life Technologies). Cells were stored at -20°C until used for cytokine ELISAs. Endotoxin concentration in OVA was determined to be 8,000 U/ml using the Limulus Amebocyte Lysate enzymatic assay QCL-1000 (Lonza Group Ltd, Basel, Switzerland) according to the manufacturer’s instructions.

**Serum and lung lavages ELISA**

To measure OVA-specific antibody titres, blood sera and lung lavages was obtained as indicated above and ELISAs were performed as previously described [33]. Immunolonal II microtiter plates (Dynex Technology Inc., Chantilly, VA, USA) were coated overnight at 4°C with OVA at 500 µg/ml in carbonate coating buffer (15 mM Na2CO3, 35 mM NaHCO3, pH 9.6; Sigma-Aldrich) and 100 µL of antigen added to each well. Wells were washed six times with distilled H2O. Pig serum or lavage samples were diluted as appropriate in Tris-buffered saline plus 0.05% Tween (Sigma-Aldrich) then they were added to the wells at 100 µL/well and incubated for two hours at room temperature (RT). Wells were washed again with distilled H2O and mouse anti-porcine IgA (Ab Serotec, Raleigh, NC, #MCA 638, 1/300), mouse anti-porcine IgG1 (Ab Serotec, #MCA 635, 1/600), mouse anti-porcine IgG2 (Ab Serotec, #MCA 636, 1/300), or mouse anti-porcine IgM (Ab Serotec, #MCA 637, 1/100) were added to the wells in a 100 µL volume and incubated for one hour at RT. Wells were washed again with dH2O and goat anti-mouse IgG (H + L) alkaline phosphatase conjugated (KPL, Gaithersburg, MD, USA, #075-1806, 1/5000) was added to each well at 100 µL/well followed by incubation for one hour at room temperature (RT). One hundred microlitres of goat anti-porcine IgG alkaline phosphatase conjugated (KPL, Gaithersburg, MD, USA, #15-14-06, 1/5000) to each well was used for total IgG detection. Wells were washed six times in dH2O before di(Tris) p-nitrophenyl phosphate (PNPP; Sigma-Aldrich) was diluted 1 mg/mL in PNPP substrate buffer (1 mM of MgCl2, 200 mM of Tris-HCl, pH 9.8; Sigma-Aldrich) and 100 µL/well was added to the wells. The reaction was allowed to develop for 60 min before absorbance was read as optical density (OD) at 405 nm in a Microplate Reader (Bio-Rad Laboratories, CA, USA). Results were reported as titres which are the reciprocal of the highest dilution that gave a positive OD reading. A positive titre was defined as an OD reading that was at least two times greater than the values for a negative sample.

**PBMC Isolation and Bioplex Cytokine Assays**

PBMCs were isolated following the protocol described by Buchanan et al [58]. Stimulation of PBMCs were performed in 96-well, round-bottom plates (Nunc, Naperville, Ill., USA) using AIM-V™ medium supplemented with 10% fetal bovine serum (Invitrogen, Burlington, ON), 2 mM L-glutamine, 50 μM 2-mercaptopetanol and 10 μg/mL polymyxin B sulfate (Sigma-Aldrich for all) as described before [12]. For each treatment, 5x10⁵ cells were cultured for 72 h in triplicate wells with media alone or 20 μg/mL OVA in 200 μL total volume. Culture supernatants were harvested and stored at -20°C until assayed for IL-10 and IFN-γ using Bioplex Cytokine Assay.

Bioplex bead coupling was performed as per the manufacturer’s instructions. The reagents were as follows: Coating antibody: monoclonal anti-swine IL-10 (Invitrogen ASC0104), Detection antibody: monoclonal anti-swine IL-10 biotin (Invitrogen ASC9109), Standard: recombinant swine IL-10 (Invitrogen PSE104) and Coating antibody: monoclonal anti-swine IFN-γ (Fisher Scientific Ltd, ENMP700), Detection antibody: monoclonal anti-swine IFN-γ (Fisher Scientific Ltd,ENMPP700; biotinylated in-house), Standard: recombinant swine IFN-γ (Ceiba Geigy). The multiplex assay was carried out in a 96 well Grenier Bio-One Flutrac 200 96 F black (VWR CanLab Mississauga, ON, #82050-754) which allows washing and retention of the LumineX beads. The porcine IFN-γ and porcine IL-10 protein standards were added to the wells at 50 µl per well at an initial concentration of 2000 pg/mL and 5000 pg/mL,
respectively followed by two-fold dilutions to make a standard curve. The PBMC supernatants were prediluted 1:3 and added to the wells at 50 μL per well. The two beadsets conjugated with the IFN-γ and IL-10 antigens were vortexed for 30 s followed by sonication for another 30 s to ensure total bead dispersal. The bead density was adjusted to 1200 beads per μl in PBS-BN (1x PBS + 1% BSA (Sigma-Aldrich) + 0.05% sodium azide (Sigma-Aldrich), pH 7.4) and 1 μL of each beadset was added to 49 μl of the PBSA + 1% New Zealand Pig Serum (Sigma-Aldrich P3484) + 0.05% sodium azide (Sigma-Aldrich) which was then added to each well. The plate was sealed with plate sealer (Fisher Scientific Ltd, #12565491) and covered with foil lid. The plate was agitated at 800 rpm for one hour at room temperature. After one hour incubation with serum, the plate was washed using the Bio-Plex ProII Wash Station (Bio-Rad Laboratories; soak 20 s, wash with 150 μL PBST three times). A 50 μL cocktail consisting of porcine IFN-γ (Fisher Scientific Ltd, (Endogen); 1/300 (which was biotinylated in-house) and biotinylated porcine IL-10 (Invitrogen; 0.5 μg/mL) was added to each well. The plate was again sealed, covered and agitated at 800 rpm for 30 minutes at room temperature then washed again as indicated above. A 50 μL volume of Streptavidin R-Phycocerythrin (RPE) (diluted to 5 μg/mL; Cedarlane, Burlington, ON; # PJRS20) was added to each well. The plate was again sealed, covered and agitated at 800 rpm for 30 minutes at room temperature and washed as indicated above. A 100 μL volume of 1 × Tris-EDTA was added to each well and then the plate was vortexed for 5 minutes before reading on the Luminex100 xMAP™ instrument (Luminex, Toronto, ON) following the manufacturer’s instructions and as described in [59]. The instrument was set up to read beadsets in regions 43 and 28 for IFN-γ and IL-10, respectively. A minimum of 60 events per beadset were read and the median value obtained for each reaction event per beadset. For all samples the multiplex assay mean fluorescent intensity (MFI) data was corrected for subtracting the background levels.

Statistical analysis

The outcome data from this study were not normally distributed and therefore, differences among experimental groups were tested using Kruskal-Wallis analysis and medians were compared using Dunn’s test. In Figure 4, significance was determined using Mann-Whitney test. Differences were considered significant if p < 0.05. All statistical analyses and graphing were formed using GraphPad Prism 5 software (GraphPad Software, San Diego, CA).

**Figure 4** OVA-specific cytokine production by PBMCs from piglets gavaged with OVA subunit vaccine then i.p. immunized with OVA/IFA at four weeks of age. Piglets (n = 5/group) were gavaged and i.p. immunized as described in Figure 1. IL-10 production was measured from PBMCs obtained four weeks post i.p. immunization. PBMCs were re-stimulated with OVA or media ex vivo. After 72 hours, the supernatants were collected and measured by BioPlex assay. Each data point represents an individual animal and median values are indicated by horizontal lines.
Abbreviations
BALS: Bronchoalveolar lavage; CMI: Cell-mediated immune response; DCs: Dendritic cells; PNPP: di(Tris) p-nitrophenyl phosphate; ELISA: Enzyme-linked immunosorbent assays; GALT: Gut-associated lymphoid tissues; h: hour; IL: Interleukin; IFA: Incomplete Freund's Adjuvant; ip: intraperitoneal; Ig: Immunoglobulin; ML: Mesenteric lymph node; MP: Microparticle; OVA: Ovalbumin; OD: Optical density; PBMC: Peripheral blood mononuclear cell; RPE: R-Phycocerythrin; RT: Room temperature; SigA: Secretory IgA; SiDev: Standard deviation; Th: T-helper cell; Treg: T regulatory cell; TBST: Tris-buffered saline with Tween 20; VIDO: Vaccine & Infectious Disease Organization.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
JAP performed the immunizations and coordinated the animal experiments; HLW conceived of and designed the experiments. JAP and HLW jointly wrote the manuscript. SN, RMB and SM performed the laboratory experiments. GKM and VG were responsible for the design of the particular vaccine component. All authors read and approved the final version.

Acknowledgments
We gratefully acknowledge financial support from the Alberta Livestock and Meat Agency, Ontario Pork and the Saskatchewan Agriculture Development Fund (Saskatchewan Ministry of Agriculture and the Canada-Saskatchewan Growing Forward bi-lateral agreement). We would like to thank the members of the Animal Care Group at VIDO for their excellent expertise as well as Donna Dent, Dr. Andrea Ladinig and Dr. John Harding for their excellent technical assistance with the BioPlex assays. We are grateful to Merital Limited for synthesis of Cpg 2395 and Idaho National Laboratory for synthesis of PCEP. HLW is an adjunct professor in the Department of Biochemistry and the School of Public Health at the University of Saskatchewan. GKM is a professor at School of Public Health at the University of Saskatchewan. VG is a professor of Veterinary Microbiology at the University of Saskatchewan. This manuscript is published with the permission of the Director of VIDO as journal series no. 701.

Author details
1. Faria AMC, Maron R, Ficker SM, Slavin AJ, Spahn T, Weiner HL. Oral tolerance induced by continuous feeding: enhanced up-regulation of transforming growth factor-beta/interleukin-10 and suppression of experimental autoimmunity encephalomyelitis. J Autoimmun. 2003;20(2):135–45.
2. Faria AMC, Weiner HL. Oral tolerance. Immunol Rev. 2005;206(1):232–59.
3. Wannemuehler MJ, Kyono H, Babb JL, Michaeli SM, Mcghee JR. Lipopolysaccharide (Lps) Regulation of the Immune-Response - Lps Converts Germ-Free Mice to Sensitivity to Oral Tolerance Induction. J. Immunol. 1992;148(3):959–65.
4. Sudu N, Sawamura SA, Tanaka K, Aiba Y, Kubo C, Koga Y. The requirement of intestinal bacterial flora for the development of an IgE production system fully susceptible to oral tolerance induction. J Immunol. 1997;159(4):1739–45.
5. Kushwah R, Hu J. Role of dendritic cells in the induction of regulatory T cells. Cell Biosci. 2011;1:120.
6. Worbs T, Sabe U, Yan S, Hoffmann MW, Hintzen G, Bernhardt G, et al. Oral tolerance originates in the intestinal immune system and relies on antigen carriage by dendritic cells. J Exp Med. 2006;203(3):199–27.
7. Alpan O, Rudomen G, Matzinger P. The role of dendritic cells, B cells, and M cells in gut-inflamed immune responses. J Immunol. 2001;166(8):4843–52.
8. Roncarolo MG, Levingis MK, Traversari C. Differentiation of T regulatory cells by immature dendritic cells. J Exp Med. 2001;193(2):F5–9.
9. Tetzka H, Ohteki T. Regulation of intestinal homeostasis by dendritic cells. Immunol Rev. 2010;234(1):247–58.
10. Pasetti MF, Simon JK, Seetin MB, Levine MM. Immunology of gut mucosal vaccines. Immunol Rev. 2011;239(1):125–48.
11. Medina E, Guzmán CA. Modulation of immune responses following antigen administration by mucosal route. FEMS Immunol Med Microbiol. 2002;47(4):305–11.
12. Nguyen TV, Yuan L, Azevedo MS, Jeong KL, Gonzalez AM, Safi LJ. Transfer of maternal cytokines to suckling piglets: in vivo and in vitro models with implications for immunomodulation of neonatal immunity. Vet Immunol Immunopathol. 2007;117(3–4):236–48.
13. Lecce JG, Matrone G. Porcine neonatal nutrition: the effect of diet on blood serum proteins and performance of the baby pig. J Nutr. 1960;70:13–20.
14. Nechvatalkova K, Kudilackova H, Leva L, Babickova K, Faldyna M. Transfer of humoral and cell-mediated immunity via colostrum in pigs. Vet Immunol Immunopathol. 2011;142(1–2):95–100.
15. Bandrick M, Antiæ-Nieto C, Baidoo SK, Molitor TW. Colostal antibody-mediated and cell-mediated immunity contributes to innate and antigen-specific immunity in piglets. Dev Comp Immunol. 2014;43(1):114–20.
16. Stott GH, Marx DB, Menefee BE, Nightengale GT. Colostal immunoglobulin transfer in calves I. Period of absorption. J Dairy Sci. 1979;62(10):1632–8.
17. Jensen AR, Elfif J, Burin DG, Sangild PT. Development of intestinal immunoglobulin absorption and enzyme activities in neonatal pigs is diet dependent. J Nutr. 2001;131(12):2359–65.
18. Halliday R. The absorption of antibodies from immune sera by the gut of the young rat. Proc R Soc Lond B Biol Sci. 1955;143(924):406–13.
19. Appleby P, Catty D. Transmission of immunoglobulin to foetal and neonatal mice. J Reprod Immunol. 1983;5(4):203–13.
20. Wend HH, Schlimp F, Geier G, Steinwender G. Time course of spontaneous bacterial translocation from gastrointestinal tract and its relationship to intestinal microflora in conventionally reared infants. Dig Dis Sci. 2001;46(5):1120–6.
21. Vukavic T. Timing of the gut closure. J Pediatr Gastroenterol Nutr. 1984;3(5):700–3.
22. Brandzaerg P, Nilssen DE, Rognon TO, Thrane PS. Ontogeny of the mucosal immune system and IgA deficiency. Gastroenterol Clin North Am. 1991;20(3):397–439.
23. Lorenz RG, Newbery RD. Isolated lymphoid follicles can function as sites for induction of mucosal immune responses. Ann N Y Acad Sci. 2004;1029:44–57.
24. Cesta MF. Normal Structure, Function, and Histology of Mucosa-Associated Lymphoid Tissue. Tox Pathol. 2006;34(5):599–608.
25. Buchanan R, Tietland S, Wilson HL. Low dose antigen exposure for a finite period in newborn rats triggers mucosal immunity rather than tolerance in later life. PLoS ONE. 2012;7(2):e31437.
26. Buchanan R, Mertins S, Wilson H. Oral antigen exposure in extreme early life in lambs influences the magnitude of the immune response which can be generated in later life. BMC Vet Res. 2013;9:160. doi:10.1186/1746-6148-9-160.
27. Pasternak JA, Ng SH, Wilson HL. A single, low dose oral antigen exposure in newborn piglets primes mucosal immunity if administered with CpG oligodeoxynucleotides and polyphosphazene adjuvants. Vet Immunol Immunopathol. 2014;161(3–4):211–21.
28. Haverson K, Corfield G, Jones PH, Kenny M, Fowler J, Bailey M, et al. Effect of Oral Antigen and Antibody Exposure at Birth on Subsequent Immune Status. Int Arch Allergy Immun. 2009;150(2):192–204.
29. Strobel S, Ferguson A. Immune responses to feed protein antigens in mice. 3. Systemic tolerance or priming is related to age at which antigen is first encountered. Pediatr Res. 1984;18(7):588–94.
30. Miller A, Lider O, Abramsky O, Weiner HL. Orally administered myelin basic protein in neonates primes for immune responses and enhances experimental autoimmune encephalomyelitis in adult animals. Eur J Immunol. 1994;24(5):1026–32.
31. Talbouts IT, Thomas WR, Holt PG. Adjuvant costimulation during secondary antigen challenge directs qualitative aspects of oral tolerance induction, particularly during the neonatal period. J Immunol. 2004;172(4):2274–85.
32. Strobel S. Immunity induced after a feed of antigen during early life: oral tolerance v. sensitisation. Proc Nutr Soc. 2001;60(4):437–42.
33. Wilson HL, Kovacs-Nolan J, Latimer L, Buchanan R, Gomis S, Babiuk L, et al. A novel triple adjuvant formulation promotes strong, Th1-biased immune responses and significant antigen retention at the site of injection. Vaccine. 2010;28(52):6286–99.
34. Gafatpati S, Eng NF, Wilson HL, Buchanan R, Mutwiri GK, Babiau LA, et al. PCPP (poly[dil(carboxylato-phenoxy)-phosphazene]) microparticel co-encapsulating ovalbumin and CpG oligo-deoxy nucleotides are potent enhancers of antigen specific Th1 immune responses in mice. Vaccine. 2010;28(52):8306–14.

35. Awate S, Wilson HL, Lai K, Babiau LA, Mutwiri G. Activation of adjuvant core response genes by the novel adjuvant PCEP. Mol Immunol. 2012;51(3):292–303.

36. Mutwiri G, Benjamin P, Soita H, Townsend H, Yost R, Roberts B, et al. Poly[dil(copper carboxylato-phenoxy)phosphazene] (PCEP) is a potent enhancer of mixed Th1/Th2 immune responses in mice immunized with influenza virus antigens. Vaccine. 2007;25(7):1204–13.

37. Mutwiri G, Bowenock TL, Babiau LA. Microparticles for oral delivery of vaccines. Expert Opin Drug Deliv. 2005;2(5):791–806.

38. Mowat AM. The regulation of immune responses to dietary protein antigens. Immunol Today. 1987;8(3):93–8.

39. Wilson HL, Obradovic MR. Evidence for a common mucosal immune system in the pig. Mol Immunol. 2014. doi: 10.1016/j.molimm.2014.09.004.

40. McNelty TN, McClure SJ, Huntley JF. Mucosal immunity in sheep and implications for mucosal vaccine development. Sm Ruminant Res. 2008;76(1–2):83–91.

41. McGhee JR, Xu-Amano J, Miller CJ, Jackson RJ, Fujihashi K, Staats HF, et al. The common mucosal immune system: from basic principles to enteric vaccines with relevance for the female reproductive tract. Reprod Fertil Dev. 1994;6(3):369–79.

42. Mestecky J. The common mucosal immune system and current strategies for induction of immune responses in external secretions. J Clin Immunol. 1987;7(4):265–76.

43. Brandtzaeg P. Regionalized immune function of tonsils and adenoids. Immunol Today. 1999;20(8):383–4.

44. Czerkinsky C, Holmgren J. Topical immunization strategies. Mucosal Immunol. 2010;3(6):545–55.

45. Bourges D, Chevaleyre C, Wang C, Berri M, Zhang X, Nicaise L, et al. Differential expression of adhesion molecules and chemokines between nasal and small intestinal mucosae: implications for T- and slgA+ B-lymphocyte recruitment. Immunology. 2007;122(4):551–61.

46. Foss DL, Murtha MP. Mucosal immunogenicity and adjuvanticity of cholera toxin in swine. Vaccine. 1999;17(7):88–90.

47. Verdonck F, De Haauwere V, Bouchaert J, Goddeeris BM, Cox E. Fimbriae of enterotoxigenic Escherichia coli function as a mucosal carrier for a coupled heterologous antigen. J Control Release. 2005;104(2):243–58.

48. Kabir S. Cholera vaccines: the current status and problems. Rev Med Microbiol. 2005;16(3):101–16.

49. Botner A, Strandbygaard B, Sorensen KJ, Have P, Madsen KG, Madsen ES, et al. Appearance of acute PRRS-like symptoms in sow herds after vaccination on a live PRRS vaccine virus. Arch Virol. 1999;144(12):2389–412.

50. Pasternak D. The common mucosal immune system and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your next manuscript to BioMed Central and take full advantage of:

Submit your manuscript at www.biomedcentral.com/submit