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Maternal Antibiotic Treatment Protects Offspring from Diabetes Development in Nonobese Diabetic Mice by Generation of Tolerogenic APCs

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Type 1 diabetes (T1D) is a T cell–mediated autoimmune disease that involves the slow, progressive destruction of islet β cells and loss of insulin production, as a result of interaction with environmental factors, in genetically susceptible individuals. The gut microbiome is established very early in life. Commensal microbiota establish mutualism with the host and form an important part of the environment to which individuals are exposed in the gut, providing nutrients and shaping immune responses. In this study, we studied the impact of targeting most Gram-negative bacteria in the gut of NOD mice at different time points in their life, using a combination of three antibiotics—neomycin, polymyxin B, and streptomycin—on diabetes development. We found that the prenatal period is a critical time for shaping the immune tolerance in the progeny, influencing development of autoimmune diabetes. Prenatal neomycin, polymyxin B, and streptomycin treatment protected NOD mice from diabetes development through alterations in the gut microbiota, as well as induction of tolerogenic APCs, which led to reduced activation of diabetogenic CD8 T cells. Most importantly, we found that the protective effect was age dependent, and the most profound protection was found when the mice were treated before birth. This indicates the importance of the prenatal environment and early exposure to commensal bacteria in shaping the host immune system and health. The Journal of Immunology, 2015, 195: 000–000.

Materials and Methods

Mice

Female NOD/Caj mice have been maintained at Yale University for many years. BDC2.5NOD and NY8.3NOD transgenic mice were purchased from The Jackson Laboratory. The mice used in this study were kept in specific pathogen-free conditions on a 12-h dark/light cycle, in individually ventilated filter cages with autoclaved food at the Yale University animal facility. The use of the animals in this study was approved by the Yale University Institutional Animal Care and Use Committee.

Antibiotic treatment

The antibiotics NPS (Sigma-Aldrich) were added to the drinking water at a final concentration of 1 mg/ml for neomycin and streptomycin and 1600 U/ml for polymyxin B.
To investigate the period in early life when mice were most susceptible to the effects of antibiotics, we treated pregnant (plugged) NOD mice with NPS (withdrawing treatment on giving birth) and observed for diabetes development in the offspring. This group was designated as NPS/preg. In a second group, newborn mice from NPS-treated mothers were sprayed with a gut bacterial suspension from the feces of adult untreated female NOD mice, once a week for 3 wk, until the mice were weaned. This group was named NPS+NOD.

To further investigate whether NPS could also inhibit diabetes development at later time points, we compared three groups of mice. As in our first experiment, the mice in the NPS/preg group were the offspring of pregnant mice receiving antibiotics in drinking water for 3 wk from mating until delivery. Mice in the NPS/born group were the newborn mice that received antibiotic through mother’s milk (the mothers were given antibiotic water for 3 wk from the date of pups’ delivery to the date of weaning). Mice in the NPS/wean group were 3-wk-old mice receiving antibiotic water for 3 wk from weaning. After the 3-wk treatment period, antibiotics were withdrawn from the water in all groups of mice.

Reagents
All fluorochrome-conjugated mAbs were purchased from BioLegend or eBioscience. The Luminex kit for cytokine measurement was purchased from Bio-Rad, and Ab-conjugated magnetic beads for T cell and APC purification were from Qiagen and Poly Scient.

Ratio of G+ and G- fecal bacteria
Bacterial DNA from mouse feces was isolated as previously described (28). The ratio of the DNA content of G+ and G- bacteria was quantified by quantitative PCR using Q5 (Bio-Rad) using G+ and G- specific forward primers (GTF and GF) and a universal bacterial reverse primer (sequences in Supplemental Table I). Total 16S rRNA was used as a positive control, and the results were analyzed by the delta-delta threshold cycle method after normalization with 16S rRNA. Each sample was analyzed in triplicate, and the experiments were repeated twice.

16S rRNA sequencing analysis
Fecal samples were collected from offspring of four to five independent breeders, and DNA extraction was performed. 16S rRNA V4 and V5 regions of fecal bacteria were amplified from DNA samples (barcoded primer sequences in Supplemental Table I). PCR products were purified by Qia gen gel extraction kit, and DNA concentration was determined with a Nano Drop spectrophotometer (Thermo Scientific) and gel electrophoresis. Following dilution to 1 × 10^6 molecules/µl, equal volumes (10 µl) were mixed together to prepare the library, followed by a final dilution to 1 × 10^7 molecules/µl. A total of 20 µl of the library was sequenced using the GS Junior Titanium Series 454 sequencing system (Roche).

Sequencing results were analyzed with the QIIME software package (http://qiime.org). Taxonomic assignment was performed using representative sequences of each picked operational taxonomic unit. β-Diversity was calculated to compare differences between microbial communities and shown as principal coordinate analysis (PCoA).

Cell purification
APCs were purified by removing CD4+ (clone GK1.5) and CD8+ T cells (clone TIB105) using mAb hybridoma supernatants, followed by magnetic bead (conjugated with goat anti-rat IgG) separation. CD4+ T cells were purified by removing CD5+ T cells (clone TIB105), MHC class II+ cells (clone 10.2.16), and B cells (anti-mouse IgM and IgG; Qiagen) using mAb hybridoma supernatants and magnetic bead separation. CD8 T cells were purified similarly, except that anti-CD4 (clone GK1.5) was used instead of anti-CD8 mAb. Individual B cells, CD11b+ cells, and CD11c+ cells were purified by EasySep Mouse B cell enrichment kit, Mouse CD11b positive selection kit, and Mouse CD11c positive selection kit II, respectively, from StemCell Technology according to the manufacturer’s instructions. The purity of the cells was routinely >90%, analyzed by flow cytometry.

T cell proliferation assay
Spleens were harvested from mice, and RBCs were lysed to obtain splenocytes. Splenocytes were counted and plated at 1 × 10^6 cells/well in a volume of 150 µl in 96-well round-bottom plates. The splenocytes were stimulated with anti-CD3 (clone 2C11 supernatant), at different dilutions, in the presence or absence of anti-CD28 (clone 37.51 supernatant; 1:100). The cells were cultured for 72 h, and the culture supernatants were collected before adding [3H]thymidine (0.5 µCi/well). The plates were further incubated for 18–19 h, the cells were harvested, and cpm were counted.

Pured CD4+ BDC2.5 or CD8+ NY8.3 T cells (10^5/well) were co-cultured with irradiated APCs (5 × 10^5/well) from 12-wk-old NOD mice in the presence or absence of BDC2.5 mimotope peptide (RTPLWVRME) or IGRPGP66-214 peptide (VYLKTNVFL), respectively. Cell proliferation was determined by stimulation index (SI), which was calculated by the formula: [3H]thymidine incorporation (cpm) with Ag/ [3H]thymidine incorporation (cpm) without Ag.

TLR agonist stimulation
Intracellular Foxp3 was detected using a Foxp3 staining kit (eBioscience) following the manufacturer’s instructions. For intracellular cytokine staining, 10^6 cells were cultured for 4 h with 50 ng/ml PMA (Sigma-Aldrich), 500 ng/ml ionomycin (Sigma-Aldrich), and 1 µl/ml GolgiPlug (BD Biosciences), before staining with Abs against surface markers. Isotype controls were used to set the gates for flow cytometric analysis. Immune cells were isolated from 12-wk-old mice.

Bacterial transfer
Newborn pups were sprayed with a freshly prepared suspension of mouse feces (from offspring of either untreated or maternally NPS-treated female NOD mice) once a week for the first 2 wk. Starting from the third week, a freshly prepared fecal suspension was administered by gavage (containing ∼1 × 10^9 culturable bacteria) to NOD recipients (either untreated or pretreated with ampicillin [1 mg/ml], metronidazole [1 mg/ml], neomycin [1 mg/ml], and vancomycin [0.5 mg/ml] [AMNN], to deplete most bacteria in the gut). Recipient mice were monitored for diabetes by weekly screening glycosuria with Diastix (Bayer), and diabetes was confirmed by blood glucose measurement (≥250 mg/dl) (FreeStyle; Abbott).

Lymphocyte adoptive transfer
Splenocytes from NPS/preg mice were injected into NOD.scid or irradiated NOD recipient mice (10^7/mouse, i.v.). The recipient mice were monitored for diabetes as described above.

APCs from NPS/preg or control NOD mice (3 × 10^6/mouse) plus T cells from diabetic NOD mice (3 × 10^6/mouse) were injected into 5-wk-old NOD.scid mice (i.v.). The recipient mice were monitored for diabetes as described above.

Insulitis scoring
Mice were dissected at the prediabetic stage (12 wk old), and pancreata were collected. Pancreata were fixed in 10% buffered formalin and then paraffin-embedded. Tissues were sectioned and stained with H&E. Insulitis was scored under light microscopy using the following grading: 0–25, no insulitis; 25–50, insulitis affecting 25–50% of the islet; 50–75, insulitis affecting 50–75% of the islet; and 75–100, >75% islet infiltration.

Statistics
Statistical analysis was performed using GraphPad Prism software (GraphPad). Any p values <0.05 were considered significant.

Results
NPS treatment has a time-dependent impact on T1D development
To investigate the effect of prenatal environment on the offspring, we treated pregnant NOD mice with NPS. We found that diabetes development in the offspring (NPS/preg) of NPS-treated parents was significantly delayed, and the incidence was also significantly reduced compared with the offspring from untreated dams (p = 0.01; Fig. 1A). It is interesting that by introducing normal gut bacteria soon after birth via spraying pups with gut bacterial suspension from nontreated adult NOD mice (NPS+NOD), the sprayed pups lost the maternal protective effect related to NPS treatment and developed a similar incidence of diabetes to the
untreated control group, although there was an ~3-wk delay in diabetes onset ($p = 0.01$ compared with NPS/preg offspring; Fig. 1A).

We further investigated the effect of NPS treatment at different time points in the early life of female NOD mice. We compared mice from mothers treated during pregnancy (NPS/preg) with two additional groups of mice—treated just after birth for 3 wk (NPS/born) and just after weaning for 3 wk (NPS/wean)—and observed the mice for diabetes. Confirming results shown in Fig. 1A, the NPS/preg mice had the most significant decrease of diabetes incidence, whereas NPS/born mice also showed delayed and reduced diabetes onset ($p < 0.05$, Fig. 1B). The incidence of diabetes in NPS/wean mice was markedly higher compared with the mice treated during the prenatal or neonatal period, although the overall incidence was still lower (55%) than in the untreated control group (80%; Fig. 1B).

Furthermore, NPS treatment led to significant reduction of insulitis compared with untreated control mice (Fig. 1C).

**NPS antibiotics altered gut microbiota**

We next investigated the change in composition of gut microbiota induced by antibiotic treatment. As expected, NPS/preg mice had a much higher ratio of $G^+ / G^-$ compared with untreated mice (Fig. 2A), measured by qPCR, which indicated that most of the G$^-$ bacteria were depleted by NPS treatment, and therefore more G$^+$ bacteria colonized the gut. It is noteworthy that although the NPS/preg mice were treated with NPS in the prenatal period only, the high $G^+/G^-$ ratio of gut bacteria in these mice persisted even at 6 mo of age (Fig. 2A), which suggests that once established, the gut microbiome is stable.

We further performed pyrosequencing of bacterial 16S rRNA of fecal samples collected from 12-wk-old NPS-treated and untreated mice, regardless of the time of NPS treatment. PCoA revealed different clusters of gut microbiota from each group of mice shown in Fig. 2B. It is interesting that NOD mice treated with NPS at different time points had distinct profiles of gut microbiota from each other, compared with untreated control NOD mice (dots).

Although the incidence of diabetes in mice from the NPS/preg and NPS/born groups was very similar, their gut microbial communities showed very different clustering (Fig. 2B, squares in top left and upward triangles in bottom right quadrant, respectively). Likewise, the incidence of diabetes in NPS/wean mice was similar to the untreated control NOD mice, whereas their gut microbial communities were very different (Fig. 2B, rightward triangles in bottom left and dots in top right quadrant, respectively). This indicated that antibiotic treatment not only significantly shifted the gut bacterial composition, but also the timing of the treatment made a substantial difference, although these differences were not always correlated with diabetes development. The pyrosequencing results showed that a major phylum of G$^-$ bacteria, Proteobacteria, were markedly reduced in NPS-treated mice compared with untreated control mice (Fig. 2C). In addition, there were also some changes in the G$^+$ bacteria, and we found that among the most abundant G$^+$ Firmicutes, two families, Lachnospiraceae and Coriobacteriaceae, showed significantly higher abundance in NPS/preg mice than in control mice (Fig. 2D).

**Diabetes protection induced by NPS treatment could be transferred to new hosts**

To investigate if the protection was long-lasting, we adoptively transferred splenocytes from NPS/preg donors to two different recipients. NOD/scid mice have no mature T cells or B cells, which makes it a good model to test the diabetogenic function of exogenous lymphocytes. The irradiated NOD mice were treated with a sublethal dose of x-rays to temporarily reduce most of the endogenous lymphocytes at the time of the transfer, but these
endogenous lymphocytes will reconstitute after x-ray treatment. Donor splenocytes from 12-wk-old NPS/preg mice and age- and sex-matched untreated mice (nondiabetic) were used as controls. As shown in Fig. 3, diabetes development was significantly delayed and reduced in both NOD.scid (Fig. 3A) and irradiated NOD (Fig. 3B) recipient mice that were injected with splenocytes from NPS/preg mice, compared with the mice injected with splenocytes from untreated NOD mice.

We also tested the protective effect of gut microbiota by fecal transplant experiments. To avoid the effects of existing gut commensals, we first treated the pregnant NOD mice with AMNV until delivery, to mimic a relatively commensal-free prenatal environment. 90% of the gut bacteria were depleted by AMNV treatment in the dams, confirmed by CFU analysis of feces; data not shown. We also set up an untreated pregnant NOD control group. Soon after birth, the offspring from each group were given gut microbiota from diabetes-free 12-wk-old NPS/preg mice. Gut microbiota from unmanipulated nondiabetic 12-wk-old NOD mice were used as a control. Diabetes development in the offspring treated with gut microbiota from NPS/preg mice was delayed and decreased, compared with the controls (the offspring that received gut microbiota from normal adult NOD mice). This phenomenon was more clearly seen in the group of recipient mice born to untreated mothers (Fig. 3D). These results indicated that diabetes protection was transferable by altered gut microbiota if endogenous microbiota of the recipients was limited.

Effects of NPS on T cell function

To investigate which immune cells contribute to this protection from diabetes in addition to the altered gut microbiota, we isolated the lymphocytes from spleen, mesenteric lymph nodes (MLN), pancreatic lymph nodes (PLN), and Peyer’s patches to study the expression of inflammatory cytokines. There were lower frequencies of both CD4 and CD8 MLN T cells expressing intracellular IFN-γ in NPS/preg mice (Fig. 4A). Consistent with the reduced frequency of IFN-γ–producing cells, the frequency of cells expressing T-bet, a master transcription factor for IFN-γ production, was also significantly lower in MLN T cells from NPS/preg mice compared with untreated control mice. Similar results were seen in PLN (Fig. 4B) and splenic T cells. Moreover, we found a reduction in the frequency of intracellular IL-17–producing CD4+ and CD8+ T cells from NPS/preg mice (data not shown). The frequency of inflammatory cytokine-producing lymphocytes in Peyer’s patches of NPS mice was also reduced (data not shown).

We then tested the effect of the antibiotic treatment on T cell function following stimulation in vitro and found that although prenatal NPS treatment did not alter T cell proliferation (Fig. 4C),
the T cells secreted less inflammatory cytokines, IFN-γ, and IL-17 (Fig. 4D).

To investigate whether regulatory T cells (Tregs) contributed to this protection, we analyzed Foxp3+ Tregs in 12-wk-old NPS/preg mice. We found that the frequency of Foxp3+ Tregs was significantly increased in all the lymphoid tissues examined (spleen, MLN, and PLN). More importantly, the absolute number of Foxp3+ Tregs was also significantly increased in spleen, MLN, and PLN of NPS/preg mice (Fig. 4E, representing FACS plot showing the frequency of Foxp3+CD25+ Tregs in splenocytes).

**Effect of NPS treatment on APCs**

We next investigated the APCs from 12-wk-old mice. The percentages of IFN-γ-producing splenic CD11c+ and IL-12 and IL-17–producing CD11b+ cells were reduced in NPS/preg mice (Fig. 5A–D). Furthermore, the CD11b+ cells expressing the im-

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**FIGURE 3.** Diabetes development by adoptive transfer. Transfer of immune cells: total splenocytes (1 × 10⁷) from offspring of untreated NOD (control) or NPS/preg (treated during gestation) NOD mice were isolated and adoptively transferred to either NOD.scid (A) (control, n = 13; NPS/preg, n = 9) or irradiated NOD mice (B) (control, n = 16; NPS/preg, n = 7). Statistical differences were analyzed by log-rank test. Transfer of gut microbiota: the recipients are the offspring of untreated or AMNV-treated (during pregnancy) mothers as described in Materials and Methods. Total gut microbiota from either nonglucogenic 12-wk-old control NOD or NPS/preg mice were transferred to offspring of AMNV-treated mothers (C) (control, n = 9; NPS/preg, n = 15; p = 0.06) or offspring of untreated NOD mothers (D) (control, n = 9; NPS/preg, n = 11). Statistical differences were analyzed by log-rank test. *p < 0.05, **p < 0.01.

**FIGURE 4.** Functional characterization of T cells from NPS/preg mice. (A) Percentage of CD4+ and CD8+ IFN-γ-expressing T cells in MLN from NOD and NPS/preg mice. Statistical differences were analyzed by t test. (B) Percentage of CD4+ and CD8+ T-bet–expressing T cells in PLN from NOD and NPS/preg groups. Statistical differences were analyzed by t test. (C) T cell proliferation in response to anti-CD3 stimulation. Splenocytes from NPS/preg mice and untreated control NOD mice were stimulated with anti-CD3 at different concentrations with or without anti-CD28. T cell proliferation was determined by [³H]thymidine incorporation and calculated as SI (cpm in the presence of anti-CD3 + anti-CD28/cpm in the absence of Abs). (D) IL-17 and IFN-γ secretion from the culture supernatants in (C). Statistical differences were analyzed by t test. (E) Representative flow cytometric plots of Tregs from splenocytes (SPL) of untreated NOD (control) or NPS/preg mice. The total number of CD25+Foxp3+ cells in gated CD4+ T cells from splenocytes, MLN, and PLN cells is shown. n = 4/group. Statistical differences were analyzed by t test. *p < 0.05, **p < 0.01, ****p < 0.0001.
munoregulatory cytokine IL-10 were increased, with a significantly higher frequency in the NPS/preg group (Fig. 5E). This strongly suggested that APCs may contribute to diabetes protection by NPS treatment.

It is known that TLRs are expressed mostly on APCs. Therefore, to test the effects of NPS treatment on the function of APCs, we stimulated splenocytes from NPS/preg mice with LPS, which is derived from the cell walls of G^- bacteria, and PGN, the major component of G^+ bacteria. Our results showed that APCs from NPS/preg mice had reduced responses to LPS but mildly enhanced responses to PGN stimulation (Fig. 6A, 6B). We further tested Ag-presenting function, using irradiated splenocytes as APCs to present Ag and stimulate the proliferation of BDC2.5 CD4^+ and NY8.3 CD8^+ diabetogenic T cells. It is interesting that NPS treatment did not affect Ag presentation and stimulation of CD4^+ T cells (Fig. 6C), but Ag presentation and stimulation of CD8^+ T cells was significantly impaired (Fig. 6D).

**FIGURE 5.** Cytokine expression profiles of APCs. (A) Representative FACS plot of IFN-γ+ cells gated on CD11c+ cells from control NOD or NPS/preg mice. (B) The percentage of IFN-γ+CD11c+ cells in splenocytes from control NOD or NPS/preg mice (n = 4/group). Statistical analysis was performed by t test. The percentage of CD11b^+IL-12+ (C), CD11b^+IL-17+ (D), and CD11b^+IL-10+ (E) cells between control NOD and NPS/preg (n = 4/group). Statistical analysis was performed by t test. *p < 0.05, **p < 0.01, ***p < 0.001.

**FIGURE 6.** Functional study of APC. (A) LPS stimulation: splenocytes from control NOD or NPS/preg mice were cultured in the presence of LPS at different concentrations for 3 d. The proliferation was determined by [3H]thymidine incorporation and presented as SI (n = 4/group). (B) PGN stimulation: the same proliferation assay was performed in the presence of PGN (n = 4/group). (C and D) Inlet autoantigen presentation assays. Purified APCs (5 × 10^4/well) from splenocytes of control NOD (n = 7) or NPS/preg (n = 6) mice were cultured with purified BDC2.5 CD4^+ T cells (1 × 10^5/well) in the presence or absence of BDC2.5 mimotope peptide for 72 h. Ag-presenting function was determined by activation and proliferation of BDC2.5 CD4^+ T cells (C). In a separate set of experiments, purified NY8.3 CD8^+ T cells (1 × 10^5/well) were tested in the presence or absence of IGRP_206-214 peptide. Ag-presenting function was determined by CD8^+ T cell proliferation to IGRP_206-214 peptide presented by APCs from control NOD or NPS/preg mice (D). Statistical differences were analyzed by two-way ANOVA. *p < 0.05, ****p < 0.0001.
To further dissect which APC subset was responsible for the reduced Ag presentation to CD8+ T cells, we purified B cells, dendritic cells (DCs), and macrophages (Mφs) from the spleens of NPS/preg donors and control NOD mice and tested their Ag-presenting function to NY8.3 CD8+ T cells. NPS treatment suppressed the Ag-presenting function of all three major APCs (Fig. 7A–C). This alteration in APC function could be mediated by the reduced production of IL-12, IFN-γ, and IL-17 but increased production of IL-10 from the APCs, shown earlier.

**APC function in vivo**

To confirm that the protection from T1D development by NPS treatment was mediated by APCs, we isolated total splenic APCs (T cell–depleted splenocytes) either from NPS/preg or control NOD mice and cotransferred these APCs with diabetogenic (T cell–depleted splenocytes) either from NPS/preg or control NOD mice, and purified total splenic APCs were isolated either from control NOD or NPS/preg mice (n = 9/group). Irradiated B cells (5 × 10^9), DCs (5 × 10^9), and Mφs (5 × 10^9) were incubated with purified CD8+ T cells (1 × 10^7) from NY8.3 NOD mice, together with different concentrations of IGRP206–214 peptide as described in Fig. 6. Statistical differences were analyzed by t test. (D) In vivo: purified total splenic APCs were isolated either from control NOD or NPS/preg mice, and purified total splenic T cells were isolated from diabetic NOD mice. Three million APCs and 2 million T cells were mixed and transferred to 5-wk-old NOD.scid mice (n = 9/group). Diabetes development was monitored twice a week by glycosuria and confirmed by blood glucose ≥250 mg/dl. Statistical differences were analyzed by log-rank test.

Is the type of antibiotic important? Recent studies have demonstrated that colonization with specific bacteria in the gut can protect mice from developing T1D. These bacteria include segmented filamentous bacteria (24), Lactobacillus johnsonii N6.2 (25), as well as Streptococcal extracts (26), and glycoprotein extracts from Klebsiella pneumoniae (27). Most of these are G+ bacteria. Concordant with these findings, in our laboratory, NOD mice treated with vancomycin, which depletes most G+ bacteria, have accelerated T1D development (Y. Hu, P. Jin, J. Peng, F.S. Wong, and L. Wen, unpublished observations). However, a recent study reported that vancomycin treatment propagated G− Akkermansia muciniphila and reduced diabetes development in NOD mice (23). Previous studies have not shown the influence of G− bacteria on diabetes development. Therefore, to investigate the effects of altering the balance of G+ and G− bacteria in the gut, we treated the NOD mice with a mixture of antibiotics to target mainly G− bacteria. Polymyxin B is selectively toxic to G− bacteria due to specificity for the LPS that exists within many G− outer membranes (28). Streptomyacin and neomycin are protein synthesis inhibitors that are antibiotics with broader spectrum but have excellent activity against G− bacteria and partial activity against G+ bacteria (29, 30). Our current study supported our hypothesis that G+/G− balance is very important to diabetes development. NPS mixture treatment would skew the gut microbial composition away from diabetogenic bacteria, and NPS resistant bacteria in the gut may have a modulatory effect on immune cells including APCs and Tregs.

Is the timing of antibiotic administration important? In human study, researchers have found distinguishing differences in the gene expression patterns in utero in infants born in contrasting standards of living and hygiene (31). This suggests the environmental factors can shape the immunity from neonatal stage and play a role in the susceptibility or protection toward autoimmune disease. Brugman et al. (32) found that BB rats treated from the time of weaning, with sulphamethoxazole, trimethoprim, and colistin sulfate, which decrease both G− and G+ bacteria, had reduced insulitis and delayed diabetes onset. Our study suggests that timing of exposure

**FIGURE 7.** Tolerogenic function of APC in vitro and in vivo. In vitro: splenic B cells (A), DCs (B), and Mφs (C) were purified from control NOD or NPS/preg mice (n = 4/group). Irradiated B cells (5 × 10^9), DCs (5 × 10^9), and Mφs (5 × 10^9) were incubated with purified CD8+ T cells (1 × 10^7) from NY8.3 NOD mice, together with different concentrations of IGRP206–214 peptide as described in Fig. 6. Statistical differences were analyzed by t test. (D) In vivo: purified total splenic APCs were isolated either from control NOD or NPS/preg mice, and purified total splenic T cells were isolated from diabetic NOD mice. Three million APCs and 2 million T cells were mixed and transferred to 5-wk-old NOD.scid mice (n = 9/group). Diabetes development was monitored twice a week by glycosuria and confirmed by blood glucose ≥250 mg/dl. Statistical differences were analyzed by log-rank test.

*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.
to antibiotics is very important. In our model system, the strongest effects were seen when antibiotics were administered in the prenatal period. It is interesting to note that diabetes protection declined when treatment was started later and suggests that the critical time window is during prenatal and neonatal periods. Our results also support the study by Hansen et al. (23), using different antibiotics, who found a significant reduction in diabetes development when NOD mice were treated with vancomycin from birth until weaning. However, when 8-week-old mice were treated, diabetes reduction became less obvious (23). In focusing on the prenatal period, Tormo-Badia et al. (33) also treated pregnant female mice with a mixture of metronidazole, neomycin, and polymyxin and found an increased incidence of diabetes in the offspring, although in their study, the treatment did not affect the time of diabetes onset. Unfortunately, the authors did not investigate other time point(s), and thus, the effect of the treatment cannot be compared with diabetes incidence at other times in their colony (33). Another recent study in pregnant females showed that a gluten-free diet reduced the incidence of diabetes in the offspring (34), but again, this was not directly compared with effects of treatment later in life (34). In our study, we sought to systematically test treatment in the three early stages of life: during pregnancy, from birth to weaning, and after weaning. Our results showed that increasing the G⁺ to G⁻ ratio reduced diabetes incidence when antibiotics were given during pregnancy and in the neonatal period, and the bacterial composition was altered. Furthermore, this pattern of bacterial colonization was persistent, confirming the importance of bacterial flora establishment in early life, as has been shown in humans (35). The treatment increased the G⁺ Firmicutes, Lachnospiraceae, and Coriobacteriaceae, and the bacterial change associated with protection from diabetes concords with the observation of a decrease in Firmicutes to Bacteroidetes ratio found in children who developed diabetes compared with healthy control subjects (36, 37). That the bacterial composition played a role in the reduction of diabetes was evident when bacterial transfer induced a delay and reduction in the onset of diabetes in the recipient mice.

How does the alteration in bacteria induce protection from diabetes? Lau et al. (25) reported that introducing Lactobacillus johnsonii strain N6.2 to BBPD rats resulted in diabetes protection, mediated by enhanced Th17 cells. We found that changing the composition of gut microbiota affected the immune system at multiple levels. These included induction of tolerogenic APCs, with reduced Ag presentation to pathogenic CD8⁺ T cells, an increase in Tregs, and an overall reduction of the inflammatory cytokine milieu. This was seen not only in the gut-associated lymphoid tissue but also in other lymphoid organs. These changes are likely to have been effected in the period when the immune system is more sensitive to environmental changes. It emphasizes the importance of early events in shaping of immune responses and determining onset of autoimmunity. Importantly, the immunotolerance induced by NPS treatment was transferrable to naive hosts, and intriguingly, APCs were the major contributor to this transferable protection.

These results have important implications for considerations of the environmental effects on the development of autoimmunity. Our study emphasizes the importance of the composition of gut microbiota that is established from birth. There is an increase in diabetes in those born by Caesarean section (38). The composition of gut bacteria is dependent on the mode of delivery, with gut microbiota in babies born by vaginal delivery more closely resembling vaginal flora but in those born by Cesarean section, the microbiota are similar to skin flora (39). Our study highlights another way that gut microbiota established at birth may be altered to a more protective type of flora and that changing the G⁺ to G⁻ bacterial balance early in life was beneficial in protecting mice from diabetes. It is likely that although individual strains may play important roles, because of heterogeneity and the gut microbe ecosystem, no single strain will be identified as being universally protective. However, discovering any such bacteria will allow further study, to increase understanding of gut microbiota interaction with the host immune system. Furthermore, identifying how bacterial metabolites may affect differentiation of the immune cells, especially APCs, and contribute to maintenance of homeostasis and self-tolerance will be particularly important.

In summary, we have made three significant findings. Firstly, targeting most of the G⁻ bacteria and some G⁺ bacteria by NPS treatment early in life significantly protected the offspring from diabetes, which indicates that early-life antibiotic usage has a long-lasting effect. Secondly, the antibiotic effect is time dependent, with better protection from diabetes development, if gut microbiota is altered from birth. Lastly, we showed that changing the composition of gut microbiota induced tolerogenic APCs, which not only failed to activate diabetogenic CD8⁺ T cells but also transferred diabetic protection to naive NOD.scid and irradiated NOD recipients. It is not yet clear how removal of G⁻ gut bacteria induced tolerogenic APCs; however, this finding opens a new area of research that may also benefit other autoimmune diseases. Ultimately, we hope to develop probiotic treatment to provide simple and effective measures of promoting symbiosis of the gut microbiome with the host and maintaining a healthy inner environment.

Disclosures
The authors have no financial conflicts of interest.

References


