

1 Gestational pertussis vaccination and the infant's cellular immune response against
2 whole-cell pertussis vaccine in the first year of life

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15 Running Head: Infant's pertussis cellular immune response

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25 **Abstract**

26 Pertussis resurgence worldwide calls for new prevention strategies, as the recently
 27 incorporated vaccine booster dose during pregnancy, whose aim is to protect newborns
 28 from infection. In Brazil, maternal Tdap vaccination is recommended since 2014, and
 29 we reported that this strategy promotes high transplacental transfer of anti-PT IgG and it
 30 is effective in protecting infants early in life. Young children are the most susceptible
 31 group and with higher mortality rates, however, it is not well known whether the
 32 elicited anti-pertussis maternal antibodies could influence in the children's immune
 33 responses further in life, especially after their own vaccination against pertussis.
 34 Considering this scenario, we conducted a study with children born to mothers who
 35 either received or not the booster dose during pregnancy, after their primary pertussis
 36 vaccination, in order to investigate the first impact of maternal immunisation on the
 37 response to infant immunisation regarding the cellular immune response, while
 38 comparing with data from the literature.

39 As transfer of maternal antibodies could result in attenuation of the immune response to
 40 vaccination in infants, this study performed to determine whether higher levels of
 41 maternal antibodies could influence in the immune response of infants to the whole-
 42 pertussis vaccination series. Results showed no difference in cytokine production
 43 between groups, a first suggestion that maternal vaccination may not interfere with
 44 recognition and cellular response generation to vaccination. This data, together with
 45 humoral immunity and epidemiological studies, is important for the implementation of
 46 maternal immunisation strategies nationwide and will contribute to assure public
 47 policies regarding vaccination schemes.

48

49 **Importance:** Pertussis, or whooping cough, is a respiratory infectious disease caused by
 50 a bacterial agent, resulting in violent coughs and possibly death in vulnerable groups,
 51 such as young children and neonates. It is known that pregnant mothers transfer
 52 antibodies to their developing foetuses for protection in early life, however anti-
 53 pertussis antibodies are not highly detected in young children. Thus, a pertussis
 54 maternal vaccination was implemented to increase maternal anti-pertussis antibodies
 55 levels in pregnant women and therefore the transference to the foetus. However,
 56 maternal antibodies can also interfere in the child immune response in the first months
 57 of life. The significance of our research is in analysing the cellular immune response of
 58 children born from pertussis-vaccinated mothers, which will give a first glimpse on how
 59 maternal antibodies could modulate the child's response to pertussis in early life.

60

61 **Introduction**

62 Pertussis is a contagious respiratory disease caused by the *Bordetella pertussis* bacteria
 63 [1]. Despite high vaccination coverage, it remains an important public health problem,
 64 re-emerging in several countries every several years [2,3]. It has a high rate of
 65 morbimortality in young children, and estimates from the World Health Organization
 66 suggest that ~50 million pertussis cases and 300,000 deaths occurred annually, mostly
 67 in children under five years of age [4]. In Brazil, between 2010 and 2014, over 22,000
 68 people were infected [5], most cases in children younger than 1 year of age [5,6].

69 The current childhood vaccination schedule includes three doses of the diphtheria,
 70 tetanus and whole cell pertussis vaccine, combined with *Haemophilus influenza* b and
 71 hepatitis B (DTwP-Hib-HBV) in the first year of life, at 8, 16 and 24 weeks of age,
 72 leaving neonates without specific protection. So, a dose of tetanus, reduced diphtheria
 73 and acellular pertussis vaccine (Tdap) during pregnancy was proposed to address this

immunity gap in young infants and promote a higher transplacental transfer of maternal antipertussis antibodies (MatAb) to the foetus, resulting in improved protection during the neonatal period and until their own vaccination scheme is completed [7].

Since 2012, this has been recommended in the United States, United Kingdom and Australia [8], and since November 2014 in Brazil [9]. Recently, we have shown Tdap maternal vaccination promotes high titers of anti-PT IgG in newborn [10], also this strategy showed a vaccine effectiveness of 82.6% in protecting infants aged <8 weeks from pertussis in Brazil [11]. However, until now, little is known whether it may affect the subsequent childhood vaccination, which uses whole-cell pertussis (wP) [12], while other countries recommend the acellular pertussis (aP) version [13]. There is also a concern that in addition to promoting protection to young infants, high MatAb titres could interfere or attenuate the immune response to the primary childhood pertussis vaccination series [14–16].

The humoral response is not the only responsible for the protection against infection, as well as there are no correlates of protection for serological levels of antipertussis antibodies [17]. Several studies show that the cellular immune response is required for effective clearance of infection from the respiratory tract and disease prevention, through effector mechanisms mediated by IFN- γ and IL-2, which are mostly produced by the T helper lymphocyte 1 (Th1). This cell type is described as having a more inflammatory profile, opposed to Th2 cells, characterized by the production of IL-4, IL-5 and IL-13. Some authors also relate the presence of IL-17 in the mechanism against *B. pertussis in vivo*, a cytokine produced by Th17 cells [18–23].

In light of this information, we sought to analyse the production of effector cytokines that could imply the activation of the infants' cellular immune response against *B.*

98 *pertussis* antigen *in vitro*, in the context of the presence of maternal anti-pertussis
99 antibodies.

100

101 **Material and Methods**

102 *Study Design and Participants*

103 This study included 43 children around 7 months of age, born either to mothers
104 vaccinated during pregnancy with a Tdap boost vaccine during the third trimester of
105 pregnancy (n=33) or mothers who did not get vaccinated (n=10). This cohort is derived
106 from a larger cohort described elsewhere [10]. Mothers were followed up after
107 parturition until their children received the primary pertussis vaccination series,
108 composed by three DTwP-Hib-HBV doses (produced by Bio-Manguinhos/Fiocruz), at
109 approximately 2, 4 and 6 months of age. The protocol was approved by the Ethics
110 Committee from the Adolfo Lutz Institute (CAAE: 37581114.0.0000.0059).

111

112 *Sample Processing and In Vitro Stimulation*

113 5 mL of heparinized venous peripheral blood samples were collected by nurses from
114 young children around one month after the third DTwP dose.

115 Samples were diluted in culture medium (RPMI 1640 [Gibco, Massachusetts/USA]
116 supplemented with 2 g of NaHCO₃, 10 mL of nonessential amino acids [Gibco], 40 mg
117 of gentamycin and 10 mL of 200 mM L-glutamine [Sigma, Missouri/USA]). Peripheral
118 blood mononuclear cells (PBMC) were isolated using a Ficoll-Paque Plus 1440 gradient
119 (GE Healthcare, Uppsala/Sweden), according to the manufacturer's instructions. Cells
120 were washed and resuspended in culture medium, counted and rested overnight (~16 h,
121 1x10⁶ cells/mL) in 48-well flat-bottom plates at 37°C in a 5% CO₂ incubator. On the
122 following day, cells were stimulated with 2 µg/mL phytohemagglutinin (PHA) for 48 h

123 [Sigma, Missouri/USA] or 5 µg/mL of inactivated pertussis toxoid (PT) for 120 h. After
124 culture, cells and supernatant were collected and stored at -80°C for later assays. Cell
125 pellets were stabilized using RNeasy Lysis Buffer (Qiagen) and phosphate buffered saline (21.02 g
126 of Na₂HPO₄, 7.16 g of NaH₂PO₄·H₂O, 164.64 g of NaCl and reverse-osmosis H₂O for 1
127 L) until total RNA extraction.

128

129 *Cytokine Detection*

130 Frozen supernatants were thawed at room temperature, and cytokines were measured
131 using both *Cytometric Bead Array Th1/Th2/Th17* kit (BD Biosciences, New
132 Jersey/USA) and *LEGENDplex Human Th Cytokine Panel (13-plex)* kit (BioLegend,
133 California/USA), according to the manufacturers' instructions, in order to compare
134 results and analyse a larger range of cytokines. Both curve and samples were read in a
135 BD LSRFortessa flow cytometer (BD Biosciences). Minimum detection limits were
136 4.88 pg/mL (BD) and 2.4 pg/mL (BioLegend). Corresponding analyses were performed
137 with FCAP Array software v3.0 (BD Biosciences) and LEGENDplex Data Analysis
138 software (BioLegend).

139

140 *RNA Extraction and Quantitative Polymerase Chain Reaction (qPCR)*

141 RNA extraction was performed using Qiagen's RNeasy Mini kit (Qiagen,
142 Hilden/Germany), following the manufacturer's instructions. To remove any traces of
143 DNA, a DNA digestion step was performed using an RNAqueous Mini Kit (Ambion,
144 Massachusetts/USA). RNA purity and quantity were evaluated by spectrophotometry in
145 a Nanodrop (Thermo Scientific, Massachusetts/USA). RNA was reverse transcribed to
146 cDNA using an iScript cDNA Synthesis Kit (Bio-Rad, California/USA) according to
147 the manufacturer's instructions.

In an attempt to identify different cell types, the chosen genes were the transcription factor *RORC2* and the cytokine gene *IL17A*, representing Th17 cells, the transcription factor *GATA3* and the cytokine gene *IL4* representing Th2 cells, and the cytokines *IFNG* for Th1 and *IL10* for regulatory response [24]. qPCR assays were run for each sample containing cDNA as a template, specific forward (5'-3') and reverse (5'-3') primers, respectively:

RORC2: TGG AAG TGG TGG TTAGGA/AAG GCT CGG AACAGCTCCAT;

GATA3: AGATGGCACGGGACACTACCT/TAATTCGGGTTCGCTTCCG;

IFNG:

GTTTTGGGTTCCTTGGCTGTTA/AAAAGAGTTCCATTATGCGCTACATC;

IL17A: GACTCCTGGGAAGACCTCATTGG/CTTGTCTCAGAATTTGGGCATCC;

IL4: ACAGCCTCACAGAGCAGAAGACT/TGTTCTTGGAGGCAGCAAAGA;

IL10: CAGGGCACCCAGTCTGAGAA/CACATGCGCCTTGATGTCTG;

GAPDH: GAAGGTGAAGGTCGGAGT/GAAGATGGTGATGGGATTTCCTCA;

and SYBR Green PCR Master Mix (Life Technologies, Massachusetts/USA) on an Applied Biosystems 7500 Real-Time PCR System (Applied Biosystems, California/USA). *GAPDH* gene expression was used as a control to normalize the data. Results were represented as fold change given by the equation described by Livak and Schmittgen [25].

Statistical Analysis

Descriptive and inferential statistical analyses were performed with GraphPad Prism software (version 5.0). Nonparametric Mann-Whitney, Kruskal-Wallis, Friedman, Dunn and Wilcoxon tests were used for inter- and intragroup analyses, and Spearman's

172 correlation coefficient was used for demographic and experimental data. We used
173 $\alpha=5\%$, 95% confidence interval, level of significance=5% and 2-sided tests.

174

175 **Results**

176 The study population consisted of young children born from women who received (VC)
177 or not (NVC) a Tdap boost dose during pregnancy (Table 1). Samples were randomly
178 chosen from a larger cohort described by Vaz-de-Lima et al. [10]. The groups presented
179 a similar and representative demographic profile when compared to the larger cohort.

180 In order to evaluate whether Tdap vaccination during pregnancy could induce
181 modification in the cellular immune response of children, we evaluated a few related
182 genes and cytokines that could represent the main cells in the cellular immune response,
183 herein described as Th1, Th2 and Th17.

184 Transcriptional profile (Figure 1) showed no difference between groups, though the
185 median levels of *GATA3*, *IL10* and *IFNG* were higher in children born to nonvaccinated
186 mothers. When comparing the normalized expression of both basal and stimulated
187 condition between groups, NVCs had lower levels for the basal condition concerning
188 *GATA3* ($p=0.0279$) and *IFNG* ($p=0.0290$) expression. In the stimulated condition, the
189 differences disappear, so the higher median level in the fold change analysis can
190 indicate that in order to achieve the same expression upon PT stimulation, NVCs must
191 have a greater increase in the expression of *GATA3* and *IFNG* due to the lower basal
192 expression.

193 To verify whether the results found in gene expression reflected the cytokine
194 production, cytokines from Th1, Th2 and Th17 profiles were quantified via flow
195 cytometry (Table 2). PHA stimulation was used as a positive control in a few samples
196 for the main cytokines for the 3 Th profiles, due to samples and kit availability.

197 In Table 2 it shown the total quantification from detectable all cytokines but IL-9 and
198 IL-17F, as those values were undetectable.

199 Overall, TNF, IFN- γ , IL-13 and IL-6 were the most produced cytokines in both groups,
200 with IL-6 being more produced in PT-stimulated cells derived from children born to
201 vaccinated mothers (VC). In contrast, IL-2 was higher is children whose mothers were
202 not vaccinated (NVC).

203 As both groups showed simultaneous IFN- γ and IL-13 production, which could indicate
204 a mixed Th1/Th2 response or a good response by individuals to all cytokines, we
205 evaluated the correlation between cytokines from different profiles, looking for a
206 predominant response in all individuals or whether some individuals were indistinctly
207 good responders to all T helper cell profiles (data not shown). However, no correlation
208 was found between groups or cytokines. Individual responses were different from each
209 other, making it challenging to find a correlation in a small group.

210 Regarding infants with higher cytokine levels, we analysed whether factors such as
211 Tdap dose, gestational age, route of birth, infant's sex, period of DTwP vaccination and
212 exclusive breastfeeding could influence the response deviation (data not shown), but no
213 significant correlations were found.

214

215 **Discussion**

216 This study was the first to show the cellular immune response in older children, after the
217 primary pertussis vaccination series, born to Tdap-vaccinated mothers in Brazil. Despite
218 the use of Tdap during pregnancy was implemented in Brazil and other countries in the
219 last few years and substantial data regarding antibody transfer and its influence on
220 children's humoral response to vaccination [10,26–28], little is known whether this
221 strategy could affect the infant's cellular immune response.

222 A few studies have been published recently about the influence of maternal antibodies
223 in the neonatal humoral response, as the study made by Ibrahim et al. [29]. Women that
224 did not receive Tdap during pregnancy but had detectable pertussis antibody levels
225 showed that maternal antibody titres at delivery did not blunt infant vaccine response
226 among children that receive at least 2 wP doses, and the authors observed that infant
227 antibody titres increased with increasing maternal antibody levels against all pertussis
228 antigens. A Brazilian study found that Tdap maternal immunization yielded
229 significantly higher anti-PT IgG levels in vaccinated mothers and their infants compared
230 to their unvaccinated counterparts and a strong positive correlation between the anti-PT
231 IgG levels in maternal and cord blood at delivery, but did not look for antibodies against
232 other pertussis antigens [10].

233 Lima et al. [28] showed the same correlation regarding antibodies against other bacterial
234 antigens, and showed for the first time the analysis of the cellular immune response in
235 whole cord blood cells, showing IFN- γ , IL-6, IL-10 and TNF production by neonatal
236 cells upon stimulation with the whole inactivated *B. pertussis*, and low levels of IL-2,
237 IL-4 and IL-17A.

238 In our study, we complemented the neonatal data by evaluating the cytokine
239 productions in older children, with the complete primary series of vaccination against
240 pertussis. The cytokines analysed were TNF- α and IL-2, which belong to a Th1 profile,
241 in association with IFN- γ ; IL-9 by Th9 cells, IL-4, IL-5 and IL-13, which contribute to
242 the Th2 response profile; IL-10 is secreted by T regulatory cells and IL-17A, IL-17F,
243 IL-21 and IL-22 secreted by Th17 cells. IL-6 is an inflammatory cytokine and is
244 important in the differentiation of Th17 cells [24].

245 We used purified PBMC stimulated with PT, for five days, as described in the literature
246 [30]. By using PBMCs it easier to access lymphocytes present in the samples and PT is

247 the only specific antigen of *B. pertussis* [31], so our group decided to first run the assays
248 with this antigen to have a first glimpse of what could have been happening, as well as,
249 with PT, we analysed the adaptive response directly related to this stimulation. We
250 could not determine the purity of PT used in the assays, but even with other bacterial
251 antigens present we could see cytokine production upon stimulus.

252 Despite using samples from older children, our study agrees with Lima et al. [28] in the
253 production of IFN- γ , IL-6 and TNF, which we found elevated in infants' samples, but
254 not IL-10. We also found low levels of IL-4 and IL-17A, but IL-13, which is an
255 indicative of the Th2 response and was not available in the assay the authors performed,
256 was found elevated in our samples. In all cytokines evaluated, there was one or more
257 individuals that we did not detect any production, indicating a strong variability
258 amongst human samples.

259 As our both groups presented similar characteristics, the slight difference in gene
260 expression could suggest modulation by vaccination. Even so, samples were limited,
261 and the mRNA analysed was just to have a glimpse of what cells populations may have
262 been present upon stimulus. Even though cytokine mRNA are produced earlier under
263 these experimental conditions, the presence of the genes can show a tendency to a
264 certain profile. This is seen in both the cytokine measurement and gene expression and
265 could be complemented by the analysis of *TBX21*, the gene responsible for Th1 cell
266 polarization [32].

267 When attempting to find correlation between different cytokines production, we found
268 that in both groups every individual produced high levels of at least one cytokine. In the
269 NVC group, a correlation between IFN- γ and IL-10 could point to a balance between
270 activation and regulation, while in the VC group the same correlation was found, as
271 well as between and IFN- γ and IL-13 (data not shown). This could indicate a mixed

272 Th1/Th2 profile, with IL-10 possibly balancing the production of IFN- γ and the
 273 activation of Th1 cells. In both groups, cytokine levels were variable between
 274 individuals, and thus, it is not possible to determine a predominant response pattern.
 275 According to Li *et al.* [33], Th2 cells do not prevent Th1 polarization but induce their
 276 apoptosis. In addition, they also observed that IL-10 could promote Th1 cell anergy
 277 through negative regulation of accessory molecules or IL-12 in antigen-presenting cells.
 278 Even inducing protection, maternal antibodies can also interfere with the child's vaccine
 279 response, reducing the vaccine efficiency [34], leading to a decrease in the child's
 280 antibody production via immune complex formation, antigen elimination via
 281 phagocytosis or epitope masking [35]. However, immune cellular response appears to
 282 be unaffected by these mechanisms [26]. As in Brazil children are vaccinated with
 283 DTwP, unlike many countries, we aimed to analyse the cellular immune response of
 284 vaccinated children born to either vaccinated or non vaccinated women, to confirm
 285 whether the cellular response would be intact. No difference was found between the two
 286 groups. Regardless of the different sample sizes and individual variabilities, both
 287 common factors in human vaccine studies [36–38], we could see IFN- γ and TNF- α
 288 production, indicative of the Th1 response, while only IL-13 from the Th2 profile was
 289 detected.

290 Overall, this work is the first step towards a complete scenario of maternal antibodies
 291 interference in cellular immune responses. Most pertussis studies describe humoral
 292 responses, before [39] and after the implementation of maternal vaccination [40,41],
 293 evaluating immunoglobulin levels produced by mothers and transferred to newborns, in
 294 countries using aP vaccines in children, unlike Brazil. A paper in Argentina describes
 295 the use of DTwP in children in the country; however, the study shows only the
 296 influence of maternal antibodies on the humoral responses of children before DTwP

297 vaccination [41]. Despite the limitations, we evidenced cellular immune response in the
 298 context of pertussis vaccination, which is shown to be important for protection [42]. Our
 299 results show no difference in cytokine production against a bacterial antigen between
 300 groups, regardless of maternal vaccination, and this could set the path to other works
 301 that will support public health policies regarding vaccination in pregnancy. This
 302 knowledge is essential to the enhancement of vaccine protocols in pregnant women and
 303 as a basis for physicians and healthcare managers to recommend this strategy.

304

305 **Conclusion**

306 This work shows cytokine production against a pertussis antigen in the context of
 307 maternal antibodies, and data could be strengthened by verifying these observations
 308 with more than one pertussis antigens. Also, there are ongoing studies that measured
 309 antibody responses to infant immunisation, in order to complement data and promote
 310 discussion about a potential difference in the impact of maternal immunisation of infant
 311 cellular and humoral responses.

312

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324

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480 Table 1. Information about the study groups

		With Tdap	Without Tdap
N (infants)		33	10
Sex (%)	Female	16 (49%)	7 (70%)
	Male	17 (52%)	3 (30%)
Route of birth (%)	Vaginal	14 (42%)	6 (60%)
	C-section	20 (58%)	4 (40%)
Exclusive breastfeeding, median (range), months		6.2 (1.8)	5.7 (2.3)
Mother's age at birth, mean (SD), years		29.9 (7.8)	24.3 (5.2)
Mother's gestational age at vaccination, mean (SD), weeks		29.6 (2.8)	-
Mother's gestational age at birth, mean (SD), weeks		39.1 (1.7)	39.6 (1.2)
Time gap between vaccination and birth, mean (SD), weeks		9.5 (2.8)	-

1st DTwP dose age (minimum-maximum, days)	21-92	62-70
2nd DTwP dose age (minimum-maximum, days)	99-155	122-194
3rd DTwP dose age (minimum-maximum, days)	182-232	183-267
Time gap between 3rd dose and sample collection (minimum-maximum, days)	10-127	7-129

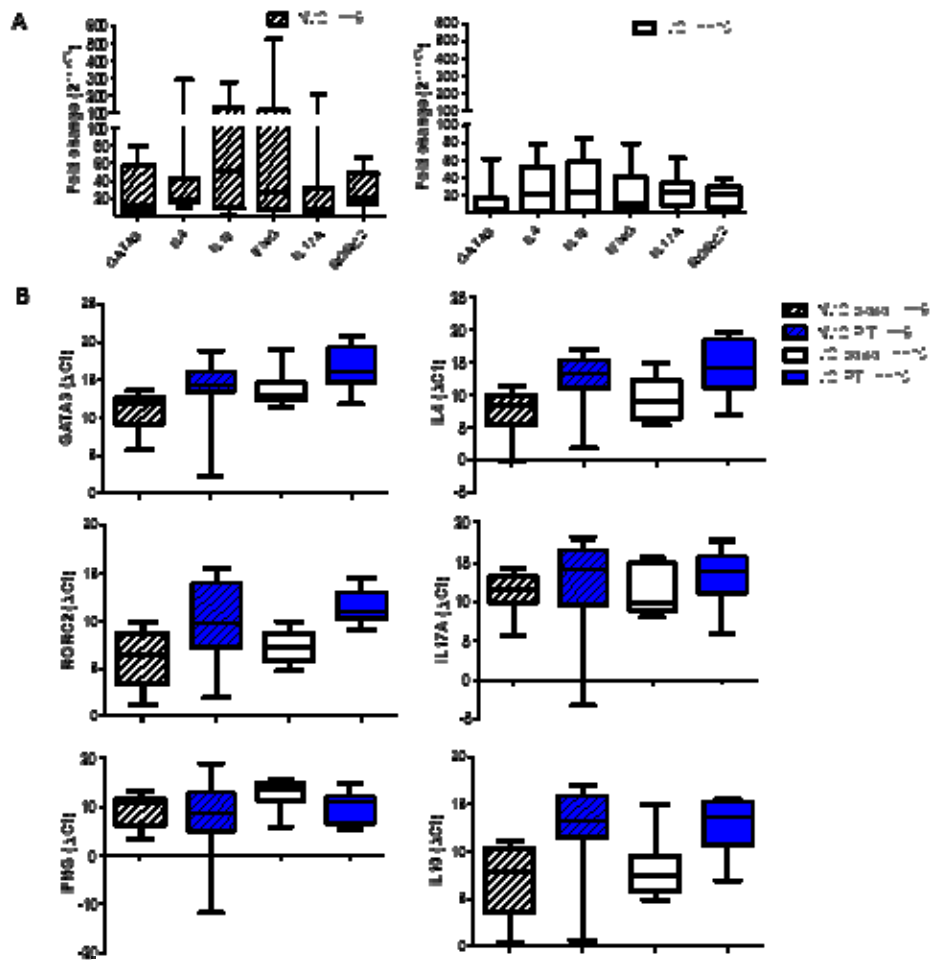
481 DTwP: diphtheria, tetanus and whole-cell pertussis vaccine; SD: standard deviation;

482 Tdap: tetanus, diphtheria and acellular pertussis vaccine.

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484 **Figure 1:** Cytokine expression in PT-stimulated PBMCs from infants born either from
485 vaccinated (VC) or nonvaccinated (NVC) mothers. (A) Relative gene expression is
486 similar in children born to Tdap-vaccinated (VC) or nonvaccinated (NVC) mothers
487 when culturing PMBCs with PT for 120 h. Box represents variation between 25th and
488 75th percentiles, and whiskers represent minimum and maximum values. (B) VC
489 presented higher basal expression of *GATA3* and *IFNG* than NVC, though differences
490 disappear with PT stimulation for 120 h. Box represents variation between the 25th and
491 75th percentiles, and whiskers represent minimum and maximum values. Statistically
492 significant differences are indicated.

493



504 Table 2. Cytokine concentration in culture supernatant of peripheral blood mononuclear cells (PBMC) from infants.

Cytokines (pg/ml)	NV				VC			
	n=2	n=2	n=10	n=10	n=4	n=4	n=33	n=28/33
	(-) basal 48h	(+) PHA	(-) basal 120h	PT	(-) basal 48h	(+) PHA	(-) basal 120h	PT
IL-5	*	*	0(0-0)	0(0-0)	*	*	0(0-0)	3,45(0- 96,54)
IL-13	*	*	0(0-0)	775,03(0- 3360)	*	*	0(0-0)	725,41(0- 5966)
IL-2	0(0-0)	102,61(15,53- 189,68)	1,3(0- 13,0)	430,25(0- 2795,84)	0,81(0-3,24)	5,45(0-12,91)	5,57(0- 94,46)	206,27-0- 4174,33)
IL-6	29,59(14,95- 44,22)	*	182,03(0- 1798)	245,24(0- 1167)	74,02(49,56- 113,11)	*	1.969,41(0- 20351)	2098,78(0- 17292)

IL-10	1,02(0,63- 1,41)	278,64(274,75- 282,54)	0,01(0- 0,1)	14,29(0- 71,73)	0,35(0-0,86)	35,42(10,83- 57,28)	2,03(0- 38,31)	21,57(0- 174,53)
IFN	0,05(0,02- 0,08)	2605,14(2058,35- 3151,94)	0,01(0- 0,1)	799,12(0- 2632,72)	0,04(0-0,08)	62,94(0,10- 248,88)	4,34(0- 127,68)	638,10(0- 3633,75)
TNF	7,42(6,58- 8,25)	2877,63(1809,63- 3945,63)	0(0-0)	303,32(0- 2292)	5,28(1,69- 8,25) N=3	370,02(105,96- 948,70)	0(0-0)	226,90(0- 2603)
IL-17A	3,73(0-7,45)	19,80(4,18- 35,41)	0,67(0- 6,68)	3,85(0- 13,86)	0(0-0)	5,94(0-23,76)	1,40(0- 8,58)	7,47(0- 32,02)
IL-4	0,85(0,58- 1,12)	3,6(1,52-5,68)	0,55(0- 1,19)	1,1(0- 2,48)	0,63(0-1,31)	1,04(0-1,9)	0,79(0- 1,45)	1,48(0- 5,5)
IL-21	*	*	0(0-0)	0(0-0)	*	*	0,66(0- 6,55)	1,045(0- 6,99)
IL-22	*	*	0(0-0)	0,954(0- 9,54)	*	*	0,18(0- 5,91)	0(0-0)

505 PBMC were stimulated *in vitro* with 2 μ L/mL PHA (+) or 5 μ g/mL pertussis toxin (PT), or left unstimulated (-) for 48 h and 120 h respectively,
506 and the supernatants collected to measure cytokine concentrations by flow cytometry. Values are presented as mean (minimum-maximum). *
507 Samples were not available for IL-5, IL-13, IL-21 and IL-22 quantification after PHA stimulation. NVC: children born from unvaccinated
508 women; VC: children born from Tdap-vaccinated women; pg/mL, picogram/milliliter.