Is there an association between socioeconomic status and
2immune response to infant and childhood vaccination in the
3Netherlands?

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13ABSTRACT

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15Introduction

16Socioeconomic status (SES) is a well-known determinant of health, but its relation with vaccine-17induced immunity is less documented. We explored the association between SES and 18immunoglobulin G (IgG) levels against vaccine-preventable diseases in vaccinated children in 19the Dutch National Immunization Programme.

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21<u>Methods</u>

22Data from a population-wide cross-sectional serosurvey in the Netherlands (2006-2007) were 23used. We compared geometric mean IgG concentrations/titers (GMC/T ratios) against measles, 24mumps, rubella, *Haemophilus influenzae* type b (Hib), *Neisseria meningococcus* type C, 25diphtheria, tetanus, poliovirus types 1,2,3 and pertussis in children of high versus low SES by 26linear regression analysis. We included 894 children (0-12 years) at one of two timeframes: 1 27month to 1 year, or 1-3 years after vaccination. Mother's educational level and net household 28income served as binary indicators of SES.

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30<u>Results</u>

31Of 58 possible associations of vaccine-induced antibody responses with educational level and 3258 with income, 10 (9%) were statistically significant: 2 favouring (that is, with higher IgG levels 33at) high educational level (for Hib 1m-1y after vaccination (GMC/T ratio: 2.99, 95%CI: 1.42-6.30)

34and polio 2 1m-1y after the 9-year booster dose (1.14, 1.01-1.27)) and 8 favouring low income 35(polio 1, 2 and 3 1m-1y after the 11-month booster (0.74, 0.58-0.94; 0.79, 0.64-0.97; 0.72, 0.55-360.95), polio 3 and pertussis 1-3y after the 11-month booster (0.70, 0.56-0.88; pertussis-prn: 370.60, 0.37-0.98; pertussis-ptx: 0.66, 0.47-0.95), mumps and rubella 1-3y after first vaccination 38(0.73, 0.55-0.97; 0.70, 0.55-0.90), and rubella 1m-1y after second vaccination (0.83, 0.55-390.90)). After adjustment for multiple testing, none of the differences remained significant. There 40was no association between SES and proportion of children with protective IgG levels.

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42Conclusion

43In this explorative study, we found no consistent association between SES and immune 44response to vaccination in the Netherlands and no association with protective IgG levels. 45Additional studies in other settings should confirm this finding.

46

48INTRODUCTION

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50Socioeconomic status (SES) is a well-known determinant of health [1]. It is a multi-faceted 51phenomenon that is at least partly captured by parameters such as education, occupational 52class and income. People with higher educational levels, from higher occupational classes and 53with higher income tend to have better health outcomes, although true causality is difficult to 54prove [2].

55The association between SES and non-communicable diseases has been studied extensively 56[3], but less is known about the effect of socioeconomic status on acute infectious diseases, 57except for its relation with the risk of exposure (e.g. crowding) and with vaccination coverage 58[4,5]. In a recent study in the Netherlands, some differences in the incidence of self-reported, 59common infectious disease syndromes (acute upper and lower respiratory tract infections, acute 60otitis media and urinary tract infections) were found between people from high versus low 61educational level, but they were not consistently in favour of either high or low educational level 62[6].

63Exposure to stress of various nature early in life has been shown to programme the immune 64system [7]. Environmental factors, including exposure to pathogens, but also psychological 65stress, poor nutrition and smoking are thought to affect one's immune response although it is 66not known to what extent [8,9]. The effect of SES on immunoglobulin-G (IgG) levels after natural 67exposure was shown to be pathogen specific and not consistently pointing to one direction in a 68recent study among adults [10]. To study the effect of SES on immune response independently 69of its association with exposure, one could compare the immune response to (childhood) 70vaccination, particularly against diseases that are no longer endemic in the area, between low 71and high SES groups. Hence, exploring the relation between SES and immune response to 72vaccination provides additional insights that could help to disentangle the complex interaction 73between SES and communicable diseases. Moreover, it might be a first step towards optimizing 74protection against vaccine preventable diseases in future. The aim of this study was to explore 75the possible association of mother's educational level and net household income (as proxy 76indicators of SES) with immune response to vaccination in infants and children vaccinated 77according to the National Immunization Programme (NIP) in the Netherlands.

78METHODS

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80Study population

81We used data from a population-wide cross-sectional serosurvey (the Pienter2 study) that was 82conducted in the Netherlands between February 2006 and June 2007. The aim of the Pienter2 83study was to establish a national serum bank to monitor antibody levels against vaccine-84preventable infectious diseases in the NIP [11]. For sampling in Pienter2, the Netherlands was 85divided in five regions and participants (0-79 years) were chosen from eight randomly selected 86municipalities in each region. People who agreed to participate, were asked to complete a 87questionnaire with questions on their background, immunisation status and health, and to 88donate a blood sample. For children younger than 14 years, a parent or guardian was asked to 89fill the questionnaire. In total, 19,781 people were invited to participate in Pienter2 which 90resulted in 6,348 (32%) completed questionnaires with supplementary blood samples, including 91an oversampling of non-Western migrants. The study was approved by the Medical Ethics 92Committee of the foundation of therapeutic evaluation of medicines (METC-STEG) in Almere 93(The Netherlands) [11].

94For our study, only children from Pienter2 who were immunized according to the NIP were 95included. At the time of the study, the NIP included DTaP-IPV-Hib (Diphtheria, Tetanus, 96Pertussis, inactivated Poliovirus and Haemophilus influenza type b) infant vaccinations at 2, 3, 4 97 and 11 months (up to 1999 at 3, 4, 5 and 11 months; Hib included since 1993), and childhood 98booster vaccinations at 4 and 9 years of age for DT-IPV (since 1962). Since 2001, the booster 99vaccination at 4 years covers pertussis as well. From 2005 onwards, the pertussis component in 100the DTP-IPV vaccine was changed from whole cell to acellular. The NIP also includes a MMR 101(measles, mumps, rubella) vaccine at the age of 14 months and 9 years. The MMR vaccine at 10214 months is combined with MenC vaccination (since 2002). Vaccination coverage at the age of 103two years was 94.3% and 94.0% in respectively 2006 and 2007 for DTP-IPV, 95.4% and 95.0% 104for Hib, 94.8% and 95.6% for MenC and 95.4% and 95.9% for MMR. At the age of 10 years, 105vaccination coverage for DT-IPV was 93.0% and 92.5% and for MMR 92.9% and 92.5% [12]. 106Dates of vaccination were copied from the vaccination booklet that participants had to bring to 107the visit where the blood sample was collected, and checked afterwards in the digital national 108immunization register. We only included children whose blood sample was taken between 1 109month and 1 year (short-term) or between 1 and 3 years (medium-term) after infant vaccination 110(that is, primary series + booster dose at 11 months of DTP-IPV and Hib; first MMR and MenC 111at 14 months), or childhood vaccination (that is, booster dose of DT-IPV or DTP-IPV at 4 years;

112booster dose of DT-IPV at 9 years; second MMR at 9 years). Furthermore, to be included in the 113study, the age range within which vaccination had to be received, was 10-14 months for the first 114booster vaccination of DTP-IPV (DTP-IPV4 scheduled at 11 months of age), 13-17 months for 115MMR1, 42-60 months for the 4-year booster vaccination of DT(P)-IPV, and 96-120 months for 116the 9-year booster of DT-IPV and MMR2. We excluded infants and children who reported to 117have been diagnosed with clinical pertussis or mumps.

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119Indicators of SES

120We used educational level of the mother and net household income as two separate indicators 121of SES, since this information was requested in the Pienter2 questionnaire. Children of whom 122no information was available on one of these indicators, were excluded from analysis with that 123indicator. To be able to include a sufficient number of children in each stratum, the indicators of 124SES were used in a binary way: low-intermediate educational level (no education, primary 125education, junior technical school, or lower general or intermediate vocational secondary 126education) versus high educational level (higher vocational or higher general secondary 127education, pre-university or university education), and low-intermediate net household income 128(≤ € 3,050/month) versus high net household income (> € 3,050/month).

129We repeated the analysis with a subset of children who belonged to the "low/intermediate-130category" for both educational level of the mother and net household income versus children 131who belonged to the "high-category" for both SES indicators to compare the extremes in a joint 132effect of educational level and household income.

133

134Laboratory analysis

135In the Pienter2 survey, IgG levels were determined by a fluorescent bead-based multiplex 136immunoassay (MMRV-MIA) using Luminex for simultaneous detection of antibodies against 137measles, mumps and rubella [13]. Antibodies against MenC and Hib were measured in a similar 138way, using combined assays [14]. Pentaplex MIA was used to determine IgG levels against 139pertussis (pertussis toxin (ptx), pertactin (prn) and filamentous hemagglutinin (FHA)), diphtheria, 140and tetanus [15]. Polio IgG total antibody levels (against poliovirus types 1, 2, and 3) were 141measured with a standard neutralization test [16]. The IgG concentrations were determined and 142calibrated to internationally accepted standards, such as the cut-off criteria of the World Health 143Organization (WHO).

144

145Data analysis

146We described the study population included after infant and childhood vaccination with 147descriptive statistics. We calculated geometric mean IgG titers/concentrations (GMC/T; with 14895% confidence intervals) for each pathogen at the two timeframes (1 month-1 year and 1-3 149years) after infant and childhood vaccination. We used linear regression analyses and 150calculated GMC/T ratios (GMC/T in the high SES groups divided by GMC/T in the low SES 151groups) to assess the effect of educational level of the mother, net household income and the 152combination of both, on logarithmically transformed IgG concentrations for the different 153pathogens at the two timeframes after vaccination. A GMC/T ratio > 1 "favoured" high 154educational level or household income (that is, antibody concentrations were higher in the high 155SES group than the low SES group). Multivariable linear regression was performed to correct 156for migration background, sex and (exact) age at vaccination. We corrected for multiple testing 157by applying the Benjamini-Hochberg's procedure on the p-values for the individual differences in 158GMC/T between low and high educational level, household income and the combination of both 159[17].

160We compared the proportions of individuals with protective levels of IgG against the different 161pathogens between children from low-intermediate (hereafter: low) and high household income, 162and between children with mothers with low-intermediate (low) and high educational level [18-16325].

164The survey design of Pienter2 with five regions (strata) and 40 municipalities (clusters) was 165taken into account in all analyses by adding them as random effects, correcting the standard 166error of the estimates. The analyses were conducted in Stata version SE/15.1 (StataCorp LLC, 167Texas, USA).

168

169Validation of results with Pienter1 data

170We repeated the analyses with data from the Pienter1 study, which was conducted between 171October 1995 and December 1996 and covered data from 8,539 participants (response rate 17256%).The Pienter1 study design was similar to Pienter2 and has been described elsewhere 173[26]. In Pienter1, only data on mother's educational level (not on household income) was 174available. At the time of the Pienter1 study, vaccination with DTP-IPV started at 3 months of age 175(3, 4, 5, and 11 months) and only the whole cell pertussis vaccine was used. MenC vaccination 176was not yet part of the NIP. Antibody levels against diphtheria and tetanus were determined 177using toxin binding inhibition assays in Pienter1; antibodies against polio by neutralization tests, 178and antibodies against measles, mumps, rubella and Hib by ELISAs [27]. For pertussis, only 179antibodies against pertussis toxin were assessed in Pienter1 by ELISA.

181**RESULTS**

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183Study population

184For the analyses by educational level, we included between 65 and 113 infants and children in 185the timeframe 1m-1y after vaccination per pathogen and between 141 and 232 infants and 186children in the timeframe 1-3y after vaccination per pathogen. For the analyses by net 187household income, these numbers were 46-101 and 117-191 respectively (Supplementary 188tables 1 and 2). Data on net household income were missing more often than data on 189educational level of the mother, which explains the difference in number of infants and children 190included. The characteristics of the study population are shown in tables 1 and 2. As expected, 191mother's educational level and net household income were correlated: infants and children of 192mothers with a low educational level were more often living in a family with a low net household 193income than infants and children of mothers with high educational level (table 1), and vice versa 194(table 2). There were significantly more children born outside the Netherlands in the low income 195and low educational level groups than in the high income and educational level groups.

197GMC/T ratios

198In figures 1 and 2, GMC/T ratios with 95% confidence intervals (CI) are presented for high 199versus low educational level of the mother and net household income respectively. A ratio >1 200means that antibody levels are higher in children with high educational level of the mother or 201with high net household income, i.e. a ratio >1 favours a high level of SES. In the analysis by 202educational level of the mother (figure 1), the GMC/T ratio (and 95% CI) was >1 for Hib 1m-1y 203after vaccination (GMC/T ratio 2.99, 95% CI 1.42-6.30) and polio 2 virus 1m-1y after the 9-year 204booster vaccination (1.14, 1.01-1.27). In the analysis by net household income (figure 2), the 205GMC/T ratio was <1 for polio 1, 2 and 3 virus 1m-1y after the 11-month booster vaccination 206(polio 1: 0.74, 0.58-0.94; polio 2: 0.79, 0.64-0.97; polio 3: 0.72, 0.55-0.95) and for polio 3 virus 207also 1-3y after the 11-month booster vaccination (0.70, 0.56-0.88). In addition, the GMC ratio 208was <1 for pertussis prn ad ptx 1-3y after the 11-month booster vaccination (prn: 0.60, 0.37-2090.98; ptx: 0.66, 0.47-0.95), for mumps 1-3 y after first vaccination (0.73, 0.55-0.97), and for 210rubella 1-3 y after first vaccination (0.70, 0.55-0.90) and 1m-1y after second vaccination (0.83, 2110.55-0.90).

212In the analysis by SES (educational level of the mother and net household income combined), 213the GMC/T ratios of rubella 1-3y after first vaccination and polio 3 virus 1-3y after the 11-month

214booster vaccination were <1 (rubella: 0.73, 0.55-0.97; polio 3: 0.68, 0.50-0.94; Supplementary 215figure 1). No other associations were found.

216After correcting for sex, migration background and age at vaccination in multivariable linear 217regression analysis, the differences in GMC/T ratio by educational level remained only 218significant for Hib, 1m-1y after vaccination(3.88; 1.97-7.66) and polio 2, 1m-1y after the 9-year 219booster (1.15; 1.02-1.30), and by net household income for polio 1 and 3, 1m-1y after the 11-220month booster dose (resp. 0.72; 0.58-0.91 and 0.73; 0.54-0.99) (Supplementary figures 2-7). In 221the multivariable regression analysis, some other differences became significant. In the 222analyses by educational level of the mother, the adjusted GMC/T ratio was 1.72 (1.07-2.76) for 223diphtheria 1m-1y after the 11-month booster vaccination, 1.36 (1.04-1.78) for tetanus and 1.23 224(1.00-1.50) for polio 2 virus 1-3y after the 11-month booster vaccination.

225After adjustment for multiple testing by applying the Benjamini-Hochberg's procedure, none of 226the differences in GMC/T between the high and low SES groups, neither in the univariable 227analyses nor in the multivariable analyses, were significant.

228

229Proportions reaching protecting IgG levels

230No differences were observed in proportions of infants and children reaching protective IgG 231levels with mothers of low versus high educational level, except for IgG levels against rubella. 232For rubella, 100% of infants of mothers with low educational level and 96% of infants of mothers 233with high educational levels reached IgG levels above the threshold for protection 1-3y after the 234first vaccination (p=0.02; table 3). In the analysis by net household income, 67% of infants from 235low income households and 50% of infants from high income households reached levels of 236protection against polio 3 virus 1-3 years after infant vaccination (p=0.04). The proportion of 237children with protective IgG levels against polio 3 is low in all children 1-3y after infant 238vaccination, but increases thereafter (table 3). This was also shown in previous studies using 239these data [28]. For the other pathogens, there were no significant differences in proportions of 240infants and children reaching protective IgG levels at the different timeframes. After adjustment 241for multiple testing, the differences between the high and low SES groups disappeared. 242

243Validation of results with Pienter1 data

244The analyses by educational level of the mother were repeated with Pienter1 data on 581 245infants aged approximately 0-4 years and 494 children 4-12 years (a total of 1,075 children). 246None of the differences in IgG levels found between children of mothers from high versus low 247educational level in the Pienter2 study were also observed in the Pienter1 study. Three 248differences were found in the Pienter1 study that were not found in Pienter2: the GMC ratios 249and 95% CI were >1 for polio 1, 2 and 3 virus 1m-1y after the 11-month booster vaccination 250(resp. 1.43 (1.03-2.01); 1.46 (1.03-2.12); and 1.57 (1.04-2.34)). These differences remained 251significant after adjusting for age and sex (data not shown), but disappeared after adjustment for 252multiple testing.

254 DISCUSSION

255In this study, we explored the effect of two indicators of SES (educational level and net 256household income) on immune response to vaccination in infants and children vaccinated 257according to the Dutch NIP. No consistent patterns were observed that favoured either high or 258low SES for any of the studied pathogens at either timeframe (1 month to 1 year after 259vaccination and 1 to 3 years after vaccination). Although a few significant differences in GMC/T 260were found for some pathogens at some timeframes, these differences were not consistent over 261timeframes, nor observed after both infant and childhood vaccination. Moreover, repetition of 262the analyses with data from the Pienter1 serosurvey that was conducted ten years earlier did 263not show similar differences but rather a few other inconsistent differences. After adjusting for 264multiple testing, all significant differences disappeared, confirming the irrelevance of the few 265differences found in the individual comparisons. The proportion of infants and children with 266protective IgG levels against the different pathogens did not differ significantly between high and 267low SES, except for slight differences for rubella and polio 3.

268Many factors may affect immune response to vaccination. Whereas there is strong evidence 269about the effect of intrinsic factors (such as age and genetics), comorbidity and vaccine factors 270on immune response to vaccination, the evidence about the relation with socioeconomic factors 271 such as nutritional status and educational level is ambiguous [9]. Studying associations 272between SES and health is complicated since several mediators and moderators along the 273 causal pathway should be considered [2]. Studies that explore the association between SES 274 and health outcomes often use educational level, income and occupation as indicators of SES, 275not in the least because they are measurable and can be addressed in policies. Whereas 276education may impact health/lifestyle behaviour, it also affects income and occupation [29]. In 277our study, low educational level was indeed associated with low net household income. 278Household income and occupation affect healthcare seeking behaviour and lifestyle, but also 279influence living conditions (e.g. crowding) and the risk of exposure to hazardous factors 280including pathogens [2, 29]. For example, several studies have shown that low SES (expressed 281in factors such as sole-parent households, maternal education, car ownership) is associated 282with increased risk of acquiring pneumococcal, Hib and meningococcal disease in the 283community [30-32].

284In a study in the Netherlands, weak associations were found between SES (educational level
285and income) and IgG concentrations induced by natural infections with rubella, measles,
286pneumococcus, Hib and MenC in non-vaccinated adults, although the direction of the
287association was not consistent (as in our study) [8]. In another study, higher IgG antibody levels

288against CMV were found in adults >25 years with lower education or income [33]. However, the 289relative contributions of differences in pathogen exposure versus differences in immune 290response after natural exposure, were difficult to assess in these studies.

291Little is known to what extent SES affects humoral immunity independently of the risk of 292exposure. By looking at the immune response after vaccination, differences in exposure can be 293ruled out, at least for vaccine-preventable diseases that are no longer prevalent in the study 294population (such as rubella, diphtheria and polio in the Netherlands). Our results do not point 295towards a clinically significant impact of SES on humoral immunity to these vaccine-preventable 296diseases in Dutch children.

297Our study had several strengths. First, we were able to use data from a national serosurvey in a 298representative sample of the Dutch population, including detailed and verified information on 299dates of vaccination for each included child [11]. Moreover; we were able to include children at 300two different timeframes after vaccination (1 month to 1 year, and 1-3 years). This allowed us to 301look at possible differences in the short versus medium-long term effects after vaccination. In 302addition, we were able to validate our results by repeating the analysis with data from the 303previous national serosurvey (10 years earlier) [26, 27].

304The study also had some limitations. As proxies for SES, only mother's educational level and 305net household income were available from the Pienter2 study. Data on possible mediators and 306moderators between these indicators and immune response to vaccination, such as nutritional 307status, smoking and alcohol use, was not collected in the Pienter2 study. Hence, even if we had 308found a clear association between education/income and immune response, we would not have 309been able to interpret this in terms of causality; additional studies with another design would be 310needed for this.

311Not for all children in the Pienter2 study, data was available on net household income. This 312resulted in smaller groups for the analysis by income and larger confidence intervals. Since 313people with a low income may be less eager to report on their income than people with higher 314incomes, the low income group may have been an underrepresentation of reality (selection 315bias). Due to small numbers in each group, we were not able to include more than two 316categories for education and net household income (low-intermediate versus high). By using 317two instead of several categories for educational level and income, we were not able to 318compare the highest versus the lowest levels of SES only, meaning that we might have missed 319differences only apparent when comparing the extremes. We compensated for this by also 320comparing GMC/T ratios in the low educational level *plus* low income group versus the high 321educational level *plus* high income group. However, in countries with relatively small differences

322in SES, such as the Netherlands, differences in immune response may be more difficult to 323detect.

324Every child was sampled only once in this cross-sectional study, meaning that every child was 325included in only one timeframe after vaccination. Thus, the two timeframes (1m-1y and 1-3y) 326could not be compared directly as data in the two timeframes were from two different groups of 327children. On the other hand, within each timeframe the data were correlated (IgG levels against 328different pathogens measured in each sample). The latter implies that an outlier in IgG level 329against one pathogen would likely be an outlier in IgG levels against other pathogens as well if a 330general factor such as SES would be the cause of this. We did not verify this at the individual 331level.

332Also, we aimed to look at immune response to vaccination only, interference with natural 333exposure to pathogens that are still circulating in the Netherlands (such as *Bordetella pertussis*, 334measles and mumps virus) could not be ruled out completely. Individuals who self-reported to 335have been diagnosed with clinical pertussis or mumps (resp. n=3 and n=0) were excluded from 336analysis, but we could not take into account possible natural boostering of immunity. In a 337previous study with Pienter2 data, an association was found between self-reported coughing > 2 338weeks in the previous 12 months and higher pertussis ptx IgG levels [34]. Although we had 339access to this information, we decided not to exclude children of whom parents reported 340coughing > 2 weeks, since that would have meant that we had to exclude about 25% of our 341study population. However, there was no difference in the numbers of infants and children with 342> 2 weeks coughing between high and low SES.

343Finally, we only considered the effect of SES on humoral immune response (IgG levels) to 344vaccination, which is still the most conventional response to investigate. However, vaccine 345response can also be quantified by looking at cellular and cytokine responses, and responses of 346the innate immune system [9]. Future studies should take this complex interplay of the different 347parts of the immune system into account.

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349In conclusion, this explorative study did not provide evidence for an association between SES 350and immune response to infant and childhood vaccination in the first three years after infant and 351childhood vaccination. Additional studies in other settings with data collected specifically for this 352purpose should confirm this. Moreover, it would be interesting to look at the longer term 353protection after vaccination in relation to SES.

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360CONFLICTS OF INTEREST

361The authors declare that they have no known competing financial interests or personal 362relationships that could have appeared to influence the work reported in this paper. 363

364CONTRIBUTORS

365JvdB, NR and MK designed the study and analysed the data. JvdB prepared the manuscript. All 366authors critically revised the manuscript. All authors approved the final article. 367

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473**TABLE 1.** Characteristics of all infants and children included in timeframes 1m-1y (n=358) and 1-3y (n=536) after vaccination

474according to NIP by educational level of mother

			Infants (approximat	tely 0-4 ye	ars)	Children (approximately 4-12 years)					
			Low edu	cational	High			Low edu	cational	High		
			level		educatio	educational		level		educational		
					level	level				level		
				%	n	%	p-value	n	%	n	%	p-value
Total			190	53%	168	47%	-	327	61%	209	39%	-
Male	e sex		92	48%	89	53%	0.27	157	48%	103	49%	0.81
Bori	<u>n in the Ne</u>	etherlands	176	93%	157	93%	0.90	269	82%	189	90%	0.002
	Indigenous	Dutch	123	65%	135	80%		199	61%	162	78%	
g	1 st generatio	on other western	0	0%	1	0.6%		1	0.3%	1	0.5%	
	2 nd generati	on other western	4	2%	11	7%		12	4%	14	7%	
kgro	1 st generatio Turkish	on Moroccan or	2	1%	1	0.6%		20	6%	2	1%	
bac	2 nd generati Turkish	2 nd generation Moroccan or Turkish		15%	1	0.6%	0.002	26	8%	1	0.5%	0.004
ion	1 st generation or Dutch Ar	on Surinam or Aruban Itillean	4	2%	2	1%		16	5%	8	4%	
grat	2 nd generation or Dutch Ar	on Surinam or Aruban Itillean	16	8%	6	4%		23	7%	8	4%	
Ξ	1 st generatio	on other non-western	2	1%	3	2%		14	4%	8	4%	
	2 nd generati	on other non-western	11	6%	8	5%		16	5%	5	2%	
Urba	anization	Very high	35	18%	31	18%		58	18%	38	18%	
		High	58	31%	54	32%	0.07	117	36%	71	34%	0.06
		Moderate	35	18%	28	17%	0.97	65	20%	45	22%	0.90
		Low	62	33%	55	33%		87	27%	55	26%	
Net		High	17	9%	73	43%		29	9%	78	37%	
hou	sehold	Low	128	67%	71	42%	<0.001	228	70%	101	48%	<0.001
inco	me	Unknown	45	24%	24	14%		70	21%	30	14%	
Med	ian age	DTP-IPV 11 m		11 (10-13)	1	1 (10-13)	0.25		n.a.		n.a.	-
(mo	nths) at	DTP-IPV 4 y		n.a.		n.a.	-	46 (44-51)		46	(44-54)	0.11
vaco	cination	DT-IPV 9 y		n.a.	n.a.		-	107 (99-116)		107 (99-114)		0.98
(5 th -9	95 th	MMR1		14 (12-16)	1	4 (13-16)	0.63	n.a.			n.a.	
(5~-95~		MMR2		n.a.		n.a.	-	107 (100-116)	107 (1	00-114)	0.96

percentile)	MenC	14 (14-16)	14 (14-16)	0.58	n.a.	n.a.	-
	Hib	11 (10-13)	11 (10-13)	0.059	n.a.	n.a.	-

477**TABLE 2.** Characteristics of all infants and children included in timeframes 1m-1y (n=294) and 1-3y (n=438) after vaccination

478according to NIP by net household income

			Infants (approximat	tely 0-4 ye	ears)	Children					
			Low hou	sehold	High hou	usehold		Low hou	sehold	High		
			income		income			income		household		
										income		
				%	n	%	p-value	n	%	n	%	p-value
Total			203	69%	91	31%	-	331	75%	107	24%	-
Male	e sex		102	50%	48	53%	0.67	155	47%	55	51%	0.47
Bori	<u>n in the Ne</u>	etherlands	183	91%	86	95%	0.42	269	81%	100	93%	0.004
	Indigenous	Dutch	122	60%	80	88%		192	58%	93	87%	
p p	1 st generatio	on other western	0	0%	1	1%	_	1	0.3%	1	0.9%	
	2 nd generati	on other western	10	5%	7	8%		16	5%	6	6%	
kgre	1 st generatio Turkish	on Moroccan or	3	1%	0	0%		18	5%	0	0%	0.007
bacl	2 nd generati Turkish	on Moroccan or	26	13%	0	0%	0.02	23	7%	0	0%	
1 st generation Surinam or Arubar or Dutch Antillean 2 nd generation Surinam or Aruba or Dutch Antillean		on Surinam or Aruban ntillean	6	3%	0	0%		18	5%	3	3%	
		on Surinam or Aruban htillean	18	9%	2	2%		25	8%	2	2%	
Σ	1 st generatio	on other non-western	6	3%	0	0%	18 20	18	5%	2	2%	
	2 nd generati	on other non-western	12	6%	1	1%		6%	0	0%		
Urba	anization	Very high	42	21%	18	20%		63	19%	21	20%	
		High	56	28%	35	38%	0 51	112	34%	43	40%	0.75
		Moderate	39	19%	13	14%	0.51	68	21%	22	21%	
		Low	66	33%	91	31%		88	27%	21	20%	
Edu	cational	High	128	63%	17	19%		101	31%	78	73%	
leve	l mother	Low	71	35%	73	80%	<0.001	228	69%	29	27%	<0.001
		Unknown	4	2%	1	1%		2	1%	0	0%	
Med	ian age	DTP-IPV 11 m		11 (10-13)	1	1 (10-13)	0.39		n.a.		n.a.	-
(mo	nths) at	DTP-IPV 4 y		n.a.		n.a.	-	4	6 (44-52)	46	(44-50)	0.55
vaco	cination	DT-IPV 9 y		n.a.	n.a.		-	107 (99-114)		107 (100-114)		0.89
(5 th -9	95 th	MMR1		14 (14-16)	1	4 (13-16)	0.81	n.a.		n.a.		-
berg	entile)	MMR2		n.a.		n.a.	-	107 (100-115)	107 (1	00-114)	0.39
percentile)		MenC		14 (14-16)	1	4 (14-16)	0.89	n.a.			n.a.	

	Hib	11 (10-13)	11 (10-13)	0.17	n.a.	n.a.	-
479							

TABLE 3. Comparison of proportions of infants and children with protective IgG levels by educational level of mother and net 482household income

			Educational level of mother					Net household income					
			Low		High			Low hou	sehold	High hou	usehold		
			educatio	nal	educatio	educational		income		income			
			level		level								
Pathogen	Threshold	Vaccination	n/N	%	n/N	%	p-value	n/N	%	n/N	%	р-	
	for											value	
	protection												
	17-24												
Measles	≥0.2 IU/ml	BMR1, 1m-1y	42/43	98%	41/41	100%	0.31	38/39	97%	27/27	100%	0.42	
		BMR1, 1-3y	126/12	100%	102/10	100%	0.28	135/13	100%	54/54	100%	-	
			6		3			5					
		BMR2, 1m-1y	62/62	100%	25/25	100%	-	56/56	100%	13/13	100%	-	
		BMR2, 1-3 y	86/86	100%	68/69	99%	0.25	88/88	100%	30/30	100%	-	
Mumps	≥45 RU/ml	BMR1, 1m-1y	40/43	93%	36/41	88%	0.42	37/39	95%	24/27	89%	0.36	
		BMR1, 1-3y	112/12	89%	88/103	85%	0.48	121/13	90%	46/54	85%	0.24	
			6					5					
		BMR2, 1m-1y	60/62	97%	25/25	100%	0.37	55/56	98%	12/13	92%	0.26	
		BMR2, 1-3 y	84/86	98%	68/69	99%	0.71	86/88	98%	30/30	100%	0.39	
Rubella	≥10 IU/ml	BMR1, 1m-1y	43/43	100%	41/41	100%	-	39/39	100%	27/27	100%	-	
		BMR1, 1-3y	126/12	100%	99/103	96%	0.02	134/13	99%	53/54	98%	0.49	
			6					5					
		BMR2, 1m-1y	61/62	98%	25/25	100%	0.54	55/56	98%	13/13	100%	0.53	
		BMR2, 1-3 y	85/86	99%	67/69	97%	0.45	87/88	99%	29/30	97%	0.41	
Diphtheria	≥0.01 IU/ml	DTP-IPV 11 m,	38/40	95%	48/48	100%	0.09	43/45	96%	28/28	100%	0.19	
		1m-1y											
		DTP-IPV 11 m, 1-	108/11	91%	78/88	89%	0.61	108/12	89%	39/44	89%	0.98	
		Зу	9					2					
		DTP-IPV 4 y, 1m-	70/70	100%	43/43	100%	-	75/75	100%	26/26	100%	-	
		1y											
		DTP-IPV 4 y, 1-3y	95/96	99%	66/67	99%	0.79	98/98	100%	35/36	97%	0.13	

			53/53	100%	21/21	100%	_	18/18	100%	12/12	100%	_
		DT-IPV 9 v 1-3v	83/83	100%	68/68	100%		87/87	100%	30/30	100%	
Tetanus	>0 01 II I/ml	DTP-IPV 11 m	40/40	100%	48/48	100%		45/45	100%	28/28	100%	
retarias	20.0110/11	1m-1v	-0/-10	10070	-0/-10	10070		-0/-0	10070	20/20	10070	
		DTP-IPV 11 m, 1-	119/11	100%	88/88	100%	-	122/12	100%	44/44	100%	-
		3y	9					2				
		DTP-IPV 4 y, 1m-	69/69	100%	43/43	100%	-	75/75	100%	26/26	100%	-
		1y										
		DTP-IPV 4 y, 1-3y	95/95	100%	67/67	100%	-	98/98	100%	35/35	100%	-
		DT-IPV 9 y, 1m-1y	53/53	100%	24/24	100%	-	48/48	100%	12/12	100%	-
		DT-IPV 9 y, 1-3y	83/83	100%	68/68	100%	-	87/87	100%	30/30	100%	-
Polio 1	Log²≥3	DTP-IPV 11 m,	39/40	98%	47/48	98%	0.90	44/45	98%	27/28	96%	0.73
		1m-1y										
		DTP-IPV 11 m, 1-	107/11	90%	82/88	93%	0.50	113/12	93%	39/44	89%	0.35
		Зу	9					2				
		DTP-IPV 4 y, 1m-	70/70	100%	43/43	100%	-	75/75	100%	26/26	100%	-
		1y										
		DTP-IPV 4 y, 1-3y	94/96	98%	66/67	99%	0.78	95/98	97%	36/36	100%	0.37
		DT-IPV 9 y, 1m-1y	53/53	100%	24/24	100%	-	48/48	100%	12/12	100%	-
		DT-IPV 9 y, 1-3y	82/83	99%	69/69	100%	0.36	87/87	100%	30/30	100%	-
Polio 2	Log²≥3	DTP-IPV 11 m,	40/40	100%	47/48	98%	0.36	45/45	100%	27/28	96%	0.21
		1m-1y										
		DTP-IPV 11 m, 1-	101/11	85%	80/88	91%	0.26	106/12	87%	39/44	89%	0.72
		Зу	9					2				
		DTP-IPV 4 y, 1m-	70/70	100%	42/43	98%	0.16	75/75	100%	25/26	96%	0.06
		1y										
		DTP-IPV 4 y, 1-3y	95/96	99%	67/67	100%	0.42	98/98	100%	35/36	97%	0.13
		DT-IPV 9 y, 1m-1y	53/53	100%	24/24	100%	-	48/48	100%	12/12	100%	-
		DT-IPV 9 y, 1-3y	82/83	99%	69/69	100%	0.36	87/87	100%	30/30	100%	-
Polio 3	Log ² ≥3	DTP-IPV 11 m,	38/40	95%	44/48	92%	0.56	44/45	98%	24/28	86%	0.06
		1m-1y										
		DTP-IPV 11 m, 1-	77/119	65%	49/88	56%	0.27	82/122	67%	22/44	50%	0.04
		Зу										
		DTP-IPV 4 y, 1m-	64/70	91%	35/43	81%	0.19	67/75	89%	22/26	85%	0.51

		1y										
		DTP-IPV 4 y, 1-3y	78/96	81%	52/67	78%	0.57	79/98	81%	28/36	78%	0.75
		DT-IPV 9 y, 1m-1y	53/53	100%	23/24	96%	0.16	48/48	100%	12/12	100%	-
		DT-IPV 9 y, 1-3y	79/83	95%	67/69	97%	0.57	85/87	98%	28/30	93%	0.28
Pertussis-	≥25 EU/ml	DTP-IPV 11 m,	24/29	83%	30/36	83%	0.95	31/35	89%	13/18	72%	0.15
prn		1m-1y										
		DTP-IPV 11 m, 1-	15/110	14%	14/85	16%	0.60	19/112	17%	5/42	12%	0.43
		Зу										
		DTP-IPV 4 y, 1m-	42/69	61%	31/45	69%	0.35	44/74	59%	19/25	76%	0.05
		1y										
		DTP-IPV 4 y, 1-3y	43/88	49%	29/53	55%	0.55	47/86	55%	14/31	45%	0.51
Hib	≥0.15 µg/ml	Hib 1m-1y	34/39	87%	46/48	96%	0.19	40/45	89%	27/28	96%	0.27
		Hib 1-3y	96/116	83%	84/93	90%	0.13	100/11	85%	43/49	88%	0.72
								7				
MenC	≥2 µg/ml	MenC 1m-1y	23/43	53%	22/42	52%	0.91	24/41	59%	14/26	54%	0.66
		MenC 1-3y	18/130	14%	11/102	11%	0.45	20/135	15%	4/56	7%	0.14
100												



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486FIGURE 1. GMC/T ratios: high versus low educational level of mother, with 95% confidence intervals



512FIGURE 2. GMC ratios: high versus intermediate/low net household income, with 95% confidence intervals

SUPPLEMENTARY TABLE 1. Number of infants and children included by educational level of mother

517SUPPLEMENTARY TABLE 2. Number of infants and children included by net household income

518SUPPLEMENTARY FIGURE 1. GMC/T ratios: high versus low SES total (net household income and educational level combined),

519 with~95% confidence intervals

SUPPLEMENTARY FIGURE 2. GMC/T ratios: high versus low educational of mother with 95% confidence intervals, unadjusted (red 522square) and adjusted (blue circle), MMR

SUPPLEMENTARY FIGURE 3. GMC/T ratios: high versus low educational of mother with 95% confidence intervals, unadjusted (red 525squares) and adjusted (red circles), Hib and MenC

SUPPLEMENTARY FIGURE 4. GMC/T ratios: high versus low educational of mother with 95% confidence intervals, unadjusted (red 528squares) and adjusted (blue circles), DTP-IPV

SUPPLEMENTARY FIGURE 5. GMC/T ratios: high versus low net household income with 95% confidence intervals, unadjusted (red 531squares) and adjusted (blue circles), MMR

SUPPLEMENTARY FIGURE 6. GMC/T ratios: high versus intermediate/low net household income with 95% confidence intervals, 534unadjusted (red squares) and adjusted (blue circles), Hib and MenC

SUPPLEMENTARY FIGURE 7. GMC/T ratios: high versus intermediate/low net household income with 95% confidence intervals, 537unadjusted (red squares) and adjusted (blue circles), DTP-IPV