ORIGINAL ARTICLE



Timeliness of childhood vaccinations in the Philippines

Peter Francis Raguindin^{1,2} · Merrylle Morales-Dizon¹ · Josephine Aldaba¹ · Lailani P. Mangulabnan³ · Renelyn P. Reyes⁴ · Batmunkh Nyambat⁵ · Maria Joyce Ducusin⁶ · Anna Lena Lopez¹

Accepted: 30 August 2020 / Published online: 4 January 2021 © World Health Organization 2020

Abstract

Timeliness of vaccinations is rarely part of monitoring in a routine immunization program. We reviewed infant immunization and conducted caregiver interviews in three regions in the Philippines from January to October 2016. We randomly selected thirty public health centers, one for each region. We defined timeliness of the receipt of antigen as within 4 weeks after the recommended age at vaccination. We assessed a total of 986 infants for timeliness of vaccination. The median age of receipt of vaccine was at 2.7 weeks (BCG), 10.1 weeks (Penta 1), and 21.7 weeks (Penta 3) compared to the recommended 0, 6, and 14 weeks of age, respectively. We found timely receipt only in 74.4% for BCG, 70.3% for Penta 1, and 39.1% for Penta 3 recipients. Thus, alongside declining immunization coverage, the infants in the Philippines had substantial delays in vaccine receipt.

Keywords Immunization \cdot Routine immunization \cdot Immunization program \cdot Immunization schedule \cdot Philippines

Introduction

Immunization is one of the most cost-effective public health interventions. According to the World Health Organization (WHO), immunization prevents between 2 and 3 million deaths every year from diphtheria, tetanus, pertussis (whooping cough), and measles [1, 2]. The WHO established the Expanded Program on Immunization (EPI) in 1974 to reduce morbidity and mortality by improving vaccine availability globally. The program initially targeted six diseases: diphtheria, pertussis, tetanus, measles, poliomyelitis, and tuberculosis [2]. Subsequently, EPI introduced newer vaccines

Electronic supplementary material The online version of this article (https://doi.org/10.1057/s4127 1-020-00255-w) contains supplementary material, which is available to authorized users.

Extended author information available on the last page of the article



 [□] Peter Francis Raguindin pnraguindin@up.edu.ph

into the program to lessen the burden of other vaccine-preventable diseases caused by hepatitis B virus, *Haemophilus influenza* type b (Hib), and *Streptococcus pneumoniae*. The EPI includes policies on evidence-based recommended vaccine schedules [2]. These schedules contain the age at which the child is most vulnerable to the disease and at which the vaccine can be given safely to achieve adequate protection.

Monitoring of immunization program performance has typically been assessed through measurements of vaccine coverage (the proportion of individuals who have received a vaccine by a benchmark age) regardless of the timing of administration [3, 4]. These do not take into account the timeliness, which detects adherence to vaccination schedules. Delays in immunization have potentially serious consequences. First, children with delayed vaccination will be unprotected from the vaccine-preventable diseases at a time when they are most at risk [5]. Second, substantial delays may contribute to diminished herd immunity or the indirect protection received by the unimmunized population when a large proportion is immunized [5]. Third, previous studies have demonstrated the association of delayed vaccination with increased risk of pertussis, measles, and *Haemophilus influenzae* type b infections and outbreaks [5–10]. Fourth, delays increase the risk of failing to achieve full immunization of the child [5, 11, 12]. Thus, adherence to immunization schedules or timeliness of vaccination is also an important indicator of a successful national immunization program aside from vaccine coverage.

In the Philippines, only one published study examined the timeliness of immunization [13]. The Philippines EPI had also been monitoring program implementation using vaccine coverage rates. In this study, we explored the timeliness of infant immunization and assess factors that may be associated with delayed vaccination using immunization records review and caregiver interviews.

Methods

Study setting and population

The Philippines is an archipelago with three major island groups (Luzon, Visayas, and Mindanao). The healthcare system is divided into 17 autonomous regional health offices charged with implementation of the immunization program. The country follows the recommended schedule by the Expanded Programme of Immunization by the World Health Organization (WHO).

We analyzed the immunization records and caregiver interviews from a study conducted on the introduction of inactivated poliovirus vaccine (IPV) in the Philippines [14]. The study collected immunization data on the following vaccines: Bacillus Calmette-Guérin vaccine (BCG) birth dose, Hepatitis B vaccine (HepB) birth dose, pentavalent vaccine series (includes vaccines for diphtheria, whole-cell pertussis, HepB, and Hib, referred to as Penta in this study), pneumococcal conjugate vaccine (13-valent) (PCV13) series, oral polio vaccine (OPV) series, and inactivated poliovirus vaccine (IPV). Because the immunization records did not contain



any sociodemographic variables about the infants, we supplemented data collection with a structured interview and investigated factors affecting vaccine timeliness.

We conducted the study from January to October 2016 in three regions, representing the three major island groups in the country: Luzon (Region 3), the Visayas (Region 6), and Mindanao (Region 10) (Fig. 1). The Philippine Department of Health selected these three regions for pilot implementation of IPV introduction. Region 3 (Central Luzon Region) has a population of 11,124,400, with an infant population (<1-year-old) of 300,300. It has 119 public health centers (PHCs) that serve the 7 provinces and 2 cities. Region 6 (Western Visayas Region) has a total population of 8,317,800, with an infant population of 224,600. It has 147 PHCs that serve the 6 provinces and 2 cities. Region 10 (Northern Mindanao Region) has a total population of 4,799,700, with an infant population of 129,600. It has 122 PHCs that serve the five provinces and 2 cities.

Field data collection

We calculated the sample size of PHCs for each region to test whether there was a 5% or greater decrease in the proportion of eligible infants receiving all the

Fig. 1 Location of the three study sites (Region 3 Central Luzon, Region 6 Western Visayas, and Region 10 Northern Mindanao)





recommended injectable vaccines with the PHCs as the primary unit of analysis. We determined 29 PHCs are needed to attain a power of 81%. We selected 30 PHCs by simple random sampling from a list in each region.

We deployed field staff already trained for interviews, data collection, and data entry in accordance with the guidelines set by the study team. We carried out additional training for interview conduct in vernacular language to maximize comprehension by interview respondents. We scheduled our staff to visit a PHC on an immunization day for data collection. An immunization day is a specific day within the week when infants received their immunizations in a PHC.

For each PHC, we prospectively reviewed the records of at least 10 infants at the site on an immunization day. If fewer than 10 infants came that day, a staff member made a return visit to comply with the minimum number. We interviewed 5 caregivers selected through convenience sampling on each immunization day in a PHC. We determined that caregivers would be eligible for interviews if≥18 years and if she or he brought an infant for that child's 14-week visit. If the study team did not achieve 5 interviews, a member of the staff had to make a return visit until having completed the minimum number. We obtained Informed consent from the caregiver. We conducted standardized interviews in an isolated place to ensure the confidentiality of the responses by the caregivers. Field staff asked the questions in vernacular language—a native dialect or form of speech of specific people or a specific region—and responses were recorded and classified for data analysis.

We collected the data using password-protected Android tablets with a preinstalled application developed for the study. The study team tabulated the answers in real time and sent them to a secured server at the study institution. We conducted data quality checks in real time to ensure accurate data entry in the application. We provided immediate feedback for any data inconsistencies and resolved these issues within the day of the visit.

The previous study focused on the introduction of IPV given at the 14th week visit and hence had data limited up to that visit only. In addition, infants may have received different vaccine dose numbers in those given in a series at that visit, depending on caregiver or healthcare provider preference or availability of vaccines during the previous visits. For example, an infant may have received Penta 1, PCV13 2, OPV 3, and IPV at this visit if s/he had not yet received these vaccines. One year after the survey, in September 2017, the study staff revisited the PHCs to gather vaccination information on the enrolled infants from the immunization records retrospectively. The PHC revisit determined whether these infants received all the recommended vaccines. We merged these data with the original dataset from the previous study to determine the timeliness of vaccination.

Definition of timely receipt and timely completion of vaccination

In this study, we used the recommended age of vaccine receipt as defined by the Philippine EPI. For infants < 6 months old, the EPI recommends these doses at birth (BCG and HepB) and visits at 6 weeks of age (Penta/PCV13/OPV dose 1), at 10 weeks (Penta/PCV13/OPV dose 2), and at 14 weeks (Penta/PCV13/OPV dose 3 and IPV). Based on this, we defined timely receipt for each vaccine, as



the receipt of the vaccine within 4 weeks after the recommended age, and we defined 'delays' as receipt of the recommended dose for > 4 weeks after the recommended age. Thus, we set age cut-off for delays at 4 weeks for BCG and HepB (recommended at birth), 10 weeks for Penta/PCV13/OPV dose 1 (recommended at 6 weeks), 14 weeks for Penta/PCV13/OPV dose 2 (recommended at 10 weeks), and 18 weeks for Penta/PCV13/OPV dose 3 and IPV (recommended at 14 weeks).

We defined 'complete vaccination' as an infant receiving all the vaccines included in the study before 6 months of age. This is an arbitrary cut-off in most immunization programs. Since maternal antibodies in an infant disappear at 6 months, most programs aim to complete the primary infant series before this age.

Data analysis

We summarized the baseline characteristics of the study population by frequency and proportion for the categorical variables. We computed for the age of receipt of vaccine for each child (expressed in weeks) by using their respective birthdate and corresponding dates of vaccination. We compared the mean age of receipt to the recommended schedule in the EPI using one-sample t test. Also, we used time-to-event analysis to determine the proportion of children having received the vaccination at each time point, as previously described in a study [15, 16]. To assess timely completion, we determined the number of infants who received all the vaccines (BCG, HepB, Penta 1, and Penta 3) on or before 6 months of age. We used logistic regression to compute the odds ratio on the timely completion and sociodemographic determinant. The sociodemographic variables included were caregiver sex (male or female), relationship to the infant, age group, number of children, and level of education. We also used the frequency of immunization days of PHCs as an additional variable in the logistic regression. As a sensitivity analysis, we performed multilevel logistic regression to determine if clustering of data had occurred because of the enrollment by regions. For statistical analysis, we used STATA 15.1 (STATA Corp, Texas, US) with two-tailed tests, and considered a P value < 0.05 significant.

Ethical consideration

Ethical clearance was obtained from the University of the Philippines Manila Research Ethics Board (UPM-REB-2015-349-01) and the WHO Regional Office of the Western Pacific Ethical Review Committee (2015.25.PHL.5.EPI) prior to the study conduct.

Results

Study population and baseline demographic data

We assessed a total of 986 infant records to determine the timeliness of vaccination, of which 465 (47.2%) had been paired with interview data



(sociodemographic factors). There was an equal distribution of infants across the different regions: 328 infants (33.3%) were from Region 3, 316 infants (32.0%) from Region 6, and 342 infants (34.7%) from Region 10. Most of the infant caregivers were females (97.8%) belonging to the 25–34 age group (46.5%) and the majority were parents of the infants (88.2). Most had at least attended secondary school (92.5%). More than half of the PHCs give routine immunizations to infants on a weekly basis (Table 1).

Table 1 Demographic data of infants and infant caregivers, and frequency of immunization in the RHUs included in the study

Characteristics	n (%)
Number of infants included	
Overall	986
Region 3	328 (33.3)
Region 6	316 (32.0)
Region 10	342 (34.7)
Infant caregiver data	
Caregivers interviewed	465 (47.2)
Sex	
Male	10 (2.2)
Female	455 (97.8)
Age	
18–24 years	159 (34.2)
25–34 years	216 (46.5)
35 years and above	90 (19.4)
Relationship to infant	
Parents	410 (88.2)
Others ^a	55 (11.8)
Number of children ^b	
0–2 children	291 (62.6)
3–4 children	122 (26.8)
5 or more children	52 (11.4)
Level of education	
Attended primary/never attended school	35 (7.5)
Attended secondary school	246 (52.9)
Attended post-secondary school	184 (39.6)
Frequency of immunization in the PHC ^c	
Weekly	301 (64.7)
2–3 times per month	30 (6.5)
Once a month	134 (28.8)

PHC public health center



^aThese include the grandparent, aunt, uncle, and other caregivers

^bOnly female caregivers were asked regarding number of children in the household

^cReflects the frequency in infants whose caregivers were interviewed

Proportion of infants vaccinated and mean age of receipt of vaccines

The vaccine coverage rates and mean age of vaccine receipt overall and per region are shown in Table 2. At the end of follow-up, all infants received Penta 1, all OPV dose series, and IPV. The proportion of vaccinated infants exceeded 95% for the rest of the vaccines except HepB, at 88%. This was true for all regions excluding Region VI, where HepB coverage exceeded 95%. The mean ages of receipt of vaccines overall were beyond the accepted range of timely receipt in the study except for BCG, HepB, and OPV 1; a similar trend was observed in all 3 regions except for Penta 1 in Regions 3 and 6. All mean values were significantly different compared to the recommended age of vaccine receipt in the Philippine EPI (Table 2).

The proportion of infants who received the vaccines at the recommended age tended to decrease with vaccine doses in a series given at a later age, as shown in the Kaplan–Meier plots (Fig. 2). At birth, only 28.1% and 62.5% of infants received BCG and HepB birth doses, with a median age of receipt of 2.7 and 0 weeks, respectively. In the primary infant series, only 3.7% of the children received Penta 1 at 6 weeks, with a median age of the receipt of 8.9 weeks of age and a delay of 2.9 weeks from the recommended age. Similarly, the proportion of children who received a timely dose of Penta 3 was smaller at 0.5%. The median age of the population that received Penta 3 was 19.7 weeks, with a delay of 5.7 weeks from the recommendation (Fig. 2a). We observed a similar trend for the other vaccine series: only 3.3% and 3.9% of infants received PCV13 1 and OPV 1 at 6 weeks, with a median delay of 3.0 and 2.4 weeks, respectively; and only 0.4% and 0.5% received PCV13 3 and IPV at 14 weeks, with a median delayed receipt of 7.7 and 5.6 weeks, respectively (Fig. 2b, c).

Timely receipt of vaccines and factors for delay in receipt of vaccines

Overall, the proportion of infants who received timely vaccination was low and below 90% for all vaccines analyzed (Tables 3). Similarly, this decreases with vaccine doses given in a series at a later age. Among the factors analyzed, a higher level of education and more frequent immunization schedule in the PHCs were associated with timely receipt of some of the vaccines. Infants of caregivers who attended secondary school were twice more likely to have infants in their care receive Penta 2 in a timely manner, while those whose caregivers attended post-secondary school were at least twice more likely to receive Penta 2, Penta 3, and OPV 3 compared to those with lower levels of education. We observed a similar trend with Penta 3 and PCV13 series among caregivers who attended secondary school, and with BCG, PCV13 2, OPV 3, and IPV among those who attended post-secondary school. Infants at PHCs that offered only a monthly immunization schedule were 40% and 50% less likely to receive BCG and HepB birth doses, respectively, compared to those with more frequent immunization schedules. On the contrary, we observed a trend towards lower chances of immunization with Penta 2, Penta 3, OPV 2, OPV 3, and IPV in infants brought to the PHCs by a female caregiver.



Table 2 Vaccination schedule and mean age of receipt among infants

)							
Vaccine	Recommended age of vaccine	Overall ^a $N = 986$		Region III ^a $N=328$		Region VI ^a $N=316$		Region X^a N = 342	
	receipt in the Philippine EPI	Proportion vaccinated at follow-up, %	Mean age at receipt, weeks (SD)	Proportion vaccinated at follow-up, %	Mean age at receipt, weeks (SD)	Proportion vaccinated at follow-up, %	Mean age at receipt, weeks (SD)	Proportion vaccinated at follow-up, %	Mean age at receipt, weeks (SD)
BCG birth dose	At birth	8.66	2.7 (4.2)	100.0	2.6 (4.1)	L'66	1.8 (3.1)	2.66	3.7 (5.0)
HepB birth dose	At birth	88.1	0.8 (3.0)	89.0	1.2 (2.7)	96.5	0.4 (3.2)	79.5	1.0 (3.2)
Penta 1	6 weeks	100.0	10.1 (5.0)	100.0	9.6 (4.4)	100.0	9.9 (5.3)	100.0	10.6 (5.3)
Penta 2	10 weeks	6.66	15.8 (6.3)	100.0	14.8 (4.7)	100.0	15.7 (6.4)	7.66	16.9 (7.2)
Penta 3	14 weeks	0.66	21.7 (7.7)	99.1	20.1 (5.6)	100.0	22.5 (8.1)	0.86	22.4 (8.6)
PCV13 1	6 weeks	99.3	10.5 (5.3)	98.5	10.0 (4.8)	100.0	10.7 (5.6)	99.4	10.7 (5.4)
PCV13 2	10 weeks	0.66	16.4 (6.6)	98.2	15.1 (4.9)	100.0	17.3 (7.6)	98.3	16.9 (6.9)
PCV133	14 weeks	87.8	22.5 (7.6)	97.9	20.2 (5.3)	99.4	24.2 (9.0)	96.2	23.1 (7.5)
OPV 1	6 weeks	100.0	9.1 (3.6)	100.0	9.0 (3.5)	100.0	8.9 (3.7)	100.0	9.4 (3.7)
OPV 2	10 weeks	100.0	14.7 (4.8)	100.0	14.2 (3.9)	100.0	14.6 (4.5)	100.0	15.4 (5.6)
OPV 3	14 weeks	100.0	20.7 (6.1)	100.0	19.4 (4.4)	100.0	21.2 (6.6)	100.0	21.4 (6.7)
IPV	14 weeks	100.0	21.0 (6.4)	100.0	19.6 (4.5)	100.0	21.6 (7.2)	100.0	21.7 (7.0)

BCG Bacillus Calmette-Guérin, HepB Hepatitis B vaccine, IPV inactivated poliovirus vaccine, OPV oral poliovirus vaccine, PCV13 pneumococcal conjugate vaccine (13-valent), SD standard deviation

^aOne sample t test using the recommended age of receipt as fixed value showed a P value of < 0.05 for all vaccines

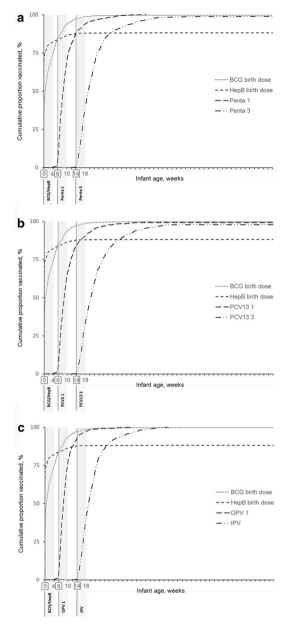


Fig. 2 Cumulative proportions of infants receiving vaccines by age in weeks. Data grouped as cumulative proportion by **a** BCG, Hepatitis B, Penta 1, and Penta 3, **b** BCG, Hepatitis B, PCV13 1, and PCV13 3, and **c** BCG, Hepatitis B, OPV 1, and IPV. Gray vertical line and boxed infant age (weeks) correspond to the recommended age of vaccine receipt. Gray area corresponds to the timely receipt as defined in this study (+ 4 weeks from recommended age). BCG, Bacillus Calmette-Guérin; HepB, Hepatitis B vaccine; IPV, inactivated poliovirus vaccine; OPV, oral poliovirus vaccine; PCV13, pneumococcal conjugate vaccine (13-valent); Penta, Pentavalent vaccine



Table 3 Factors for timely receipt^a of vaccines among infants whose caregivers were interviewed (*N*=465)

	20		Hep B DIT	ndose	Penta 1	-	Penta 2	2	Penta 3	33
	Timely	OR (P	Timely OR	OR (P	Timely	OR (P	Timely	OR (P	Timely	OR (P
	vaccination, %	value)	vaccination, %	value)	vaccination, %	value)	vaccination, %	value)	vaccination, %	
Overall	74.4		80.0		70.3		54.0		39.1	
Caregiver characteristics										
Female	97.4	:	97.6	,	97.3	:	96.4	:	96.2	1
Male	2.6	3.2 (0.279)	2.4	2.3 (0.437)	2.8	3.8 (0.201)	3.6	7.9 (0.051)	3.9	3.7 (0.059)
Relationship to infant										
Parent	88.7	;	87.4	,	88.1	;	88.1	;	89.0	1
Others	11.3	0.8 (0.527)	12.6	1.5 (0.285)	11.9	1.0 (0.919)	12.0	1.0 (0.928)	11.0	0.9 (0.653)
Age group										
18-24 years old	33.5	;	33.6	1	35.1	:	35.9	;	36.8	1
25-34 years old	46.2	1.1 (0.808)	46.5	1.1 (0.727)	45.3	0.8 (0.426)	44.6	0.8 (0.362)	44.5	0.8 (0.364)
35 years old and	20.3	1.3 (0.401)	19.9	1.3 (0.496)	19.6	0.9 (0.838)	19.5	0.9 (0.742)	18.7	0.8 (0.501
above										
Number of children										
0-2 children	63.0	;	64.5	1	62.4	;	64.1	;	66.5	1
3-4 children	25.4	0.9 (0.556)	25.0	0.7 (0.144)	26.9	1.1 (0.680)	25.1	0.9 (0.493)	23.6	0.8 (0.231)
5 or more children	11.6	1.1 (0.757)	10.5	0.6 (0.205)	10.7	(989.0) 6.0	10.8	0.9 (0.650)	6.6	0.7 (0.347
Level of education										
Attended primary school	6.7	:	7.8	1	6.4	:	4.8	;	4.4	1
Attended secondary	50.6	1.3 (0.511)	51.3	0.7 (0.485)	54.4	1.7 (0.136)	53.0	2.3 (0.032)	53.3	2.2 (0.063)
school										
Attended post-	42.8	2.1 (0.057)	40.9	1.0 (0.972)	39.1	1.5 (0.268)	42.2	2.6 (0.013)	42.3	2.4 (0.039)
secondary school										
Frequency of immunization in the PHC	in the PHC									
Weekly	69.7	;	6.99	,	64.8	;	67.7	;	65.4	1
2-3 times a month	6.1	0.5 (0.200)	8.1	1 ()	8.0	2.7 (0.069)	6.0	0.8 (0.496)	7.1	1.2 (0.686)
Once a month	24.3	(0000)	25.0	0.5 (0.002)	٠ ٢ ٢	0 8 (0 403)	26.3	0.7 (0.162)	376	177 07 0 0



Table 3 (continued)

iable 5 (collulaca	- II		1										į	
Characteristics	PCV13 1		PCV13 2		PCV13 3		OPV 1		OPV 2		OPV 3		Δd	
	Timely vaccination, %	OR (P value)												
Overall	0.99		49.3		34.4		77.4		60.2		41.9		40.7	
Caregiver characteristics	istics													
Female	07.4		96 9		96 9		97.2	;	8 90	:	96.4		26.3	
Male	2.6	2.1	3.1	2.4	3.1	1.9	2.8	1(-)	3.2	6.1	3.6	3.3	3.7	3.5
		(0.356)		(0.198)		(0.302)				(0.087)		(0.085)		(0.072)
Relationship to infant	ıt													
Parent	87.6	;	87.3	,	88.1		88.3		87.9	:	88.7	:	88.9	:
Others	12.4	1.2	12.7	1.2	11.9	1.0	11.7	6.0	12.1	1.1	11.3	6.0	11.1	6.0
Age group		(0.609)		(0.583)		(0.982)		(0.842)		(0.796)		(0.757)		(0.692)
18-24 years	36.2	;	36.7	;	35.6	;	34.7	;	35.4	;	36.9	;	35.5	1
plo														
25-34 years	44.6	0.7	43.7	0.8	45.6	6.0	45.3	0.8	44.3	0.8	43.6	0.8	45.0	6.0
plo		(0.197)		(0.211)		(0.680)		(0.475)		(0.344)		(0.250)		(0.587)
35 years old	19.2	8.0	19.7	6.0	18.8	6.0	20.0	1.1	20.4	1.0	19.5	6.0	19.6	1.0
and above		(0.488)		(0.668)		(0.689)		(0.796)		(0.867)		(0.640)		(0.875)
Number of children														
0-2 children	61.9	;	62.5	1	66.3	:	61.7	1	62.9	;	66.2	1	9.59	1
3-4 children	27.4	1.2	26.2	1.0	23.1	0.8	28.1	1.5	27.1	1.1	24.6	0.8	24.9	8.0
		(0.485)		(0.994)		(0.235)		(0.146)		(0.730)		(0.351)		(0.442)
5 or more	10.8	0.9	11.4	1.0	10.6	0.8	10.3	0.8	10.0	0.8	9.5	0.7	9.5	0.7
children		(0.799)		(606.0)		(0.605)		(0.429)		(0.370)		(0.194)		(0.282)
Level of education														
Attended	6.2	;	5.2	:	4.4	;	6.7	;	6.1	;	4.6	;	4.8	1
primary school														
Attended	55.4	1.9	54.6	2.0	56.3	2.3	53.6	1.7	53.9	1.7	53.3	2.1	53.4	2.0
secondary		(0.084)		(0.071)		(0.059)		(0.196)		(0.151)		(0.066)		(0.086)
school														
Attended post-	38.4	1.5	40.2	1.9	39.4	2.1	39.7	1.6	40.0	1.6	42.1	2.3	41.8	2.2
secondary		(0.272)		(0.091)		(0.103)		(0.247)		(0.178)		(0.042)		(0.061)
school														
Frequency of immunization in the PHC	nization in the F	HC												
Weekly	65.8	;	65.5	;	61.9	;	68.1	;	70.0	;	2.99	;	9.59	1
2-3 times a	8.9	1.1	5.7	8.0	7.5	1.4	7.5	2.1	6.1	0.7	7.2	1.2	7.4	1.2
month		(0.748)		(0.498)		(0.433)		(0.249)		(0.359)		(0.714)		(0.563)
Once a month	27.4	8.0	28.8	1.0	30.6	1.2	24.4	0.4	23.9	0.5	26.2	8.0	27.0	6.0
		(0.370)		(0.954)		(0.455)		(0.000)		(0.003)		(0.317)		(0.538)

Bold represents the P value of < 0.05 is considered statistically significant

BCG Bacillus Calmette-Guérin, HepB Hepatitis B vaccine, IPV inactivated poliovirus vaccine, OPV oral polio vaccine, OR odds ratio, Penta pentavalent vaccine, PHC public health center, IPV inactivated poliovirus vaccine, PCV13 pneumococcal conjugate vaccine (13-valent)
^aTimely receipt of vaccines is defined as receipt of vaccines within 4 weeks from recommended age based on the Philippine EPI



Table 4 Factors for timely completion^a of vaccines among infants whose caregivers were interviewed (N=465)

Caregiver characteristics	Timely completion, n (%)	Delayed completion, <i>n</i> (%)	OR (P value)
Overall	282 (60.7)	183 (39.4)	
Sex			
Female	274 (97.2)	181 (98.9)	_
Male	8 (2.8)	2 (1.1)	2.6 (0.2)
Relationship to infant			
Parent	245 (86.9)	165 (90.2)	_
Others	37 (13.1)	18 (9.8)	1.4 (0.3)
Age group			
18–24 years	93 (33.0)	66 (36.1)	_
25–34 years	127 (45.0)	89 (48.6)	1.0 (1.0)
35 years and above	62 (22.0)	28 (15.3)	1.6 (0.1)
No. of children of female caregivers			
0–2 children	185 (65.6)	106 (57.9)	_
3–4 children	71 (25.2)	51 (27.9)	0.8 (0.3)
5 or more children	26 (9.2)	26 (14.2)	0.6 (0.07)
Level of education			
Attended primary school/never attended school	119 (42.2)	65 (35.5)	_
Attended secondary school	145 (51.4)	101 (55.2)	1.4 (0.4)
Attended post-secondary school	18 (6.4)	17 (9.3)	1.7 (0.1)

BCG Bacillus Calmette-Guérin, HepB Hepatitis B vaccine, IPV inactivated poliovirus vaccine, OPV oral poliovirus vaccine, OR odds ratio, Penta pentavalent vaccine, PCV13 pneumococcal conjugate vaccine (13-valent)

^aTimely completion of vaccines is defined as 3 doses of Penta, PCV13, and OPV, and one dose each of BCG birth dose, HepB birth dose, and IPV received within 6 months of age

Because we conducted the sampling by region, we performed a multilevel logistic regression model (Supplement Table 1) as part of our sensitivity analyses. In this analysis using regions as clusters, we found no statistically relevant factors for the delay in vaccine receipt.

Timely completion of immunization and factors for delayed completion

Of the 465 infants whose caregivers we interviewed, only 60.7% completed all the vaccines on time. We observed no statistically significant difference among caregivers' age, sex, and educational status for the delayed completion of primary infant series. However, there was a trend towards delayed completion of immunization for infants in the care of females with 5 or more children in the household (Table 4).



Discussion

In most developing countries with problems on immunization coverage, information about timeliness of immunization is not routinely collected. Instead, most countries monitor receipt of EPI vaccines at 12 or 24 months. In the Philippines, the EPI monitors 'fully immunized children' (FIC), defined as infants who received BCG, 3 doses of DPT, 3 doses of OPV, 3 doses of HepB, and one dose of anti-measles vaccine before reaching 1 year of age [17]. However, this information does not provide insights into the extent to which vaccinations are administered on time. Our study revealed a substantial delay in the vaccine receipt for infants. The receipt of age-appropriate vaccination was delayed with only 60.7% having completed vaccination at 6 months of age, and the mean age of vaccine receipt more than 4 weeks later than the recommended age (Penta 1 recommended at 6 weeks but received at 10.1 weeks, Penta 2 recommended at 10 weeks but received at 15.8 weeks, and Penta 3 recommended at 14 weeks but received at 21.7 weeks).

We conducted the study after a 6- to 9-month pentavalent stockouts that occurred in 2015, and this resulted in untimely vaccination. Hence, the information on the extent of the delays may not be representative of the current state of immunization. The World Health Organization and UNICEF estimated that repeated stockouts of vaccines contributed to a 15% point reduction in the Philippine vaccine coverage for 2015, but the coverage marginally improved in the subsequent years [18]. According to the 2017 National Demographic and Health Survey in the Philippines, only 70% of children aged 12–23 months received all the recommended vaccinations [19]. With low immunization coverage and delayed vaccination, a rise in the number of vaccine-preventable diseases appeared in the following years. Following the repeated Pentavalent vaccine stockouts in the Philippines, the WHO Representative Office of the Philippines reported a rise in the number of diphtheria cases in the country in 2015 [20]. The declining immunization coverage also resulted in a measles outbreak in January 2019 [21] and the re-emergence of polio in September 2019 [22].

We identified further limitations in our study. First, our analysis is limited to primary infant immunization series (BCG, Hep B birth dose, DTwP-Hib-HepB, PCV, and OPV-IPV), which limits our analysis up to children 6 months of age. Equally important are the vaccination delays and coverage among older infants receiving measles and measles-mumps-rubella (MMR) vaccine. Studies have shown increased dropouts with increasing infant age [23, 24]. Second, we did not obtain other important maternal and social factors, namely, family income, distance from the PHCs, and other indicators for health-seeking behaviors. Third, our study has a small sample size, as we only included infants nested in a survey. A larger study to determine when children are being vaccinated in the Philippines will be useful for disease control and prevention efforts as well as for policymakers, as this will provide information in ascertaining the appropriate vaccination schedule.

Our findings confirm delays in immunization in the Philippines that are similar to other developing countries [24–28]. The combination of low immunization



coverage and the delayed vaccine receipt among fully immunized children has negative repercussions on the control of vaccine-preventable diseases. These may result in a reduction in the benefits of the immunization program if there are delays in protecting high-risk children. With the delays, the age of the susceptible population becomes younger, and further disease transmission is shifted to a much younger population.

Our results explored the timeliness of vaccination in a country with a hugely growing and young population. In 2015, the Philippines added an additional injectable vaccine, IPV, to the schedule. Concomitant multiple injectable vaccination may have been a factor in changing and/or delaying scheduled immunizations, and healthcare providers' perceptions and attitudes may have been a major contributing factor [14]. The work schedule of each PHCs is a major driving factor. In some PHCs, immunization visits are scheduled once per week up to once per month. The latter precludes an increased number of opportunities for the infant to be vaccinated at PCHs with less frequent immunization sessions. This then translates to low immunization coverage and further delays in vaccination.

Another finding of our study is the delay in the receipt of BCG and HepB birth doses. For BCG, one of the most commonly cited reasons for disparity in the delay is the health worker's striving to open a new vial only when a minimum number of vaccines/infants are reached to be cost-efficient. A BCG multidose vial is only available for 6 h after reconstitution, and the vial should be discarded if not used [1]. Thus, these providers advised caregivers to go to the nearest immunization facility for the infant to receive the vaccine in order to minimize the vaccine wasting. This practice is not allowed in the current immunization guidelines. The guideline requires health workers to open vials and vaccinate infants regardless of whether the minimum number of vaccines/infants is reached. This practice is not a problem with HepB vaccine as the multidose vial can be used within 28 days after opening, and the utility of each vial can be maximized easily. Disparities in Hepatitis B delays are brought about by difficulties in reviewing records. The vaccine is usually given in birthing facilities while the immunization registries and immunization cards are being distributed in immunization facilities [29]. PHCs serve as both birthing facilities and immunization facilities, but this practice may differ across all regions.

Conclusion

We have updated the data on the timeliness of immunization for infants in the Philippines and explored delays in vaccinations that could be common for low-middle income countries with a huge vaccine target population. With the looming threat of vaccine hesitancy that may cause more delays in vaccination or even vaccine refusals, the gains achieved against vaccine-preventable diseases are at risk [30]. Timeliness of receipt is crucial to confer early protection to an infant, to sustain herd immunity in the population, and to minimize the lost opportunity for vaccine completion. With the resurgence of vaccine-preventable diseases worldwide, we highlight the need for continued monitoring of vaccination, including delays in the



timing of vaccines. Timeliness of vaccine receipt is not only a surrogate measure for vaccine hesitancy in the population but has an impact on the control of vaccine-preventable disease in the population.

Acknowledgements We would like to acknowledge the help of Vincent Sumergido (Region 6 Western Visayas), Shiela Laquindanum (Region 3 Central Luzon), and Karen Hojas (Region 10 Northern Mindanao) for their support in the continuing data collection activities. We would like to thank Luzviminda Garcia for her coordinating work in the Department of Health Central Office and Brendalyn Red for her administrative help in the National Institutes of Health-UP Manila. We would also like to thank Abigail Marasigan for rendering the figures used in this paper.

Disclaimer The findings and conclusions in this report are those of the author(s) and do not necessarily represent the official position of the U.S. Centers for Disease Control and Prevention. The authors have no conflicts of interest.

Author contributions PFR: Conceptualization, Methodology, Analysis, Writing, Editing. MPM: Methodology, Data curation, Analysis, Writing, Editing. JA: Methodology, Analysis, Writing, Editing. LM: Resources, Supervision, Analysis, Editing. RR: Resources, Supervision, Analysis, Editing. NB: Methodology, Analysis, Writing, Reviewing Editing. ALL: Conceptualization, Methodology, Analysis, Writing, Editing, Funding acquisition.

Funding Open access funding provided by University of Bern. The study used funds from a project supported by the U.S. Centers for Disease Control and Prevention. The funding agency has no had no role in the design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and was ultimately responsible for the decision to submit this work for publication.

Compliance with ethical standards

Conflicts of interest The authors declare no conflicts of interest.

Ethical approval The study obtained clearance from the University of the Philippines Manila Research Ethics Board (UPM-REB-2015-349-01) and the WHO Regional Office for the Western Pacific Ethical Review Committee (2015.25.PHL.5.EPI).

The opinions expressed in this article are those of the authors and do not necessarily reflect the views of the Food and Agriculture Organization of the United Nations (FAO), its Board of Directors, or the countries they represent.

Open Access This article is licensed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 3.0 IGO License, which permits any non-commercial use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the Food and Agriculture Organization of the United Nations (FAO), provide a link to the Creative Commons licence, and indicate if changes were made. If you remix, transform, or build upon this article or a part thereof, you must distribute your contributions under the same licence as the original. The use of the Food and Agriculture Organization of the United Nations (FAO)'s name, and the use of the Food and Agriculture Organization of the United Nations (FAO) shall be subject to a separate written licence agreement between the Food and Agriculture Organization of the United Nations (FAO) and the user and is not authorized as part of this CC-IGO licence. Note that the link provided above includes additional terms and conditions of the licence. The images or other third party material in this article are included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by-nc-sa/3.0/igo/.



References

 World Health Organization. Immunization in practice: a practical guide for health staff. Geneva: WHO Press; 2015.

- Feikin DR, Flannery B, Hamel MJ, Stack M, Hansen PM. Vaccines for Children in Low- and Middle-Income Countries. In: Black RE, Laxminarayan R, Temmerman M, Walker N, editors. Reproductive, maternal, newborn, and child health: disease control priorities, vol. 2. 3rd ed. Washington (DC): The International Bank for Reconstruction and Development / The World Bank © 2016.
- Luman ET, Barker LE, Shaw KM, McCauley MM, Beuhler JW, Pickering LK. Timeliness of childhood caccinations in the United States: days undervaccinated and number of vaccines delayed. J Am Med Assoc. 2005;293(10):1024–211.
- Masters NB, Wagner AL, Boulton ML. Vaccination timeliness and delay in low- and middleincome countries: a systematic review of the literature, 2007-2017. Hum Vacc Immunother. 2019;15(12):2790-805.
- Guerra FA. Delays in immunization have potentially serious health consequences. Paediatr Drugs. 2007;9(3):143–8.
- Choudhary TS, Reddy NS, Apte A, Sinha B, Roy S, Nair NP, et al. Delayed vaccination and its predictors among children under 2 years in India: insights from the national family health survey-4. Vaccine. 2019;37(17):2331–9.
- Grant CC, Roberts M, Scragg R, Stewart J, Lennon D, Kivell D, et al. Delayed immunisation and risk of pertussis in infants: unmatched case-control study. BMJ. 2003;326(7394):852–3.
- Kolos V, Menzies R, McIntyre P. Higher pertussis hospitalization rates in indigenous Australian infants, and delayed vaccination. Vaccine. 2007;25(4):588–90.
- Siedler A, Hermann M, Schmitt HJ, Von Kries R. Consequences of delayed measles vaccination in Germany. Pediatr Infect Dis J. 2002;21(9):826–30.
- von Kries R, Böhm O, Windfuhr A. Haemophilus influenzae b-vaccination: the urgency for timely vaccination. Eur J Pediatr. 1997;156(4):282–7.
- Kiely M, Boulianne N, Talbot D, Ouakki M, Guay M, Landry M, et al. Impact of vaccine delays at the 2, 4, 6 and 12 month visits on incomplete vaccination status by 24 months of age in Quebec, Canada. BMC Public Health. 2018;18(1):1364.
- 12. Lieu T, Black S, Sorel M, Ray P, Shinefield H. Would better adherence to guidelines improve childhood immunization rates? Pediatrics. 1997;98:1062–8.
- 13. Wallace AS, Sobel H, Ryman TK, Mantaring JB 3rd, Silvestre M, Thorley M, et al. Timing of hepatitis B vaccination and impact of non-simultaneous vaccination with DTP vaccine following introduction of a hepatitis B birth dose in the Philippines. J Pub Health Policy. 2012;33(3):368–81.
- Lopez AL, Harris JB, Raguindin PF, Aldaba J, Morales M, Sylim P, et al. Introduction of inactivated poliovirus vaccine in the Philippines: effect on health care provider and infant caregiver attitudes and practices. Vaccine. 2018;36(48):7399–407.
- Dayan GH, Shaw KM, Baughman AL, Orellana LC, Forlenza R, Ellis A, et al. Assessment of delay in age-appropriate vaccination using survival analysis. Am J Epidemiol. 2006;163(6):561–70.
- Laubereau B, Hermann M, Schmitt HJ, Weil J, von Kries R. Detection of delayed vaccinations: a new approach to visualize vaccine uptake. Epidemiol Infect. 2002;128(2):185–92.
- Epidemiology Bureau DOH. Electronic field health service information system 2011: manual of operations. Manila: DOH; 2011. https://uhmis.doh.gov.ph/index.php/downloads/155-manual-of-opera tion/167-electronic-field-health-service-information-system-2011.
- WHO, UNICEF. WHO-UNICEF Joint report form for immunization Geneva: WHO and UNICEF;
 2017. https://data.unicef.org/wp-content/uploads/country_profiles/Philippines/Immunization_phl.pdf.
- Philippine Statistics Authority, ICF. Philippines National Demographic and Health Survey 2017: key indicators. Quezon City, Philippines: PSA and ICF; 2018.
- WHO Representative Office Philippines. Lessons learned from recent diphtheria cases: Working towards sustainable prevention and treatment of vaccine-preventable diseases Manila: WHO; 2017. http://www.wpro.who.int/philippines/mediacentre/features/lessons_learned_diphtheria_cases/en/. Accessed 31 Mar 2019.
- International Federation of Red Cross. Philippines: Re-emergence of vaccine preventable diseases— Measles outbreak. Geneva, Switzerland: International Federation of Red Cross and Red Crescent Societies; 2019. Contract No.: MDRPH032.



- International Federation of Red Cross. Philippines: Re-emergence of vaccine preventable diseases (polio). Geneva, Switzerland: International Federation of Red Cross and Red Crescent Societies; 2019. Contract No.: MDRPH032.
- 23. Akmatov MK, Mikolajczyk RT. Timeliness of childhood vaccinations in 31 low and middle-income countries. J Epidemiol Commun Health. 2012;66(7):e14.
- 24. Clark A, Sanderson C. Timing of children's vaccinations in 45 low-income and middle-income countries: an analysis of survey data. Lancet. 2009;373(9674):1543–9.
- Mbengue MAS, Mboup A, Ly ID, Faye A, Camara FBN, Thiam M, et al. Vaccination coverage and immunization timeliness among children aged 12-23 months in Senegal: a Kaplan-Meier and Cox regression analysis approach. Pan Afr Med J. 2017;27(Suppl 3):8.
- Akmatov MK, Kretzschmar M, Kramer A, Mikolajczyk RT. Timeliness of vaccination and its effects on fraction of vaccinated population. Vaccine. 2008;26(31):3805–11.
- Zaidi SM, Khowaja S, Kumar Dharma V, Khan AJ, Chandir S. Coverage, timeliness, and determinants
 of immunization completion in Pakistan: evidence from the Demographic and Health Survey (200607). Hum Vaccin Immunother. 2014;10(6):1712–20.
- 28. Gibson DG, Ochieng B, Kagucia EW, Obor D, Odhiambo F, O'Brien KL, et al. Individual level determinants for not receiving immunization, receiving immunization with delay, and being severely underimmunized among rural western Kenyan children. Vaccine. 2015;33(48):6778–85.
- 29. Sobel HL, Mantaring JB 3rd, Cuevas F, Ducusin JV, Thorley M, Hennessey KA, et al. Implementing a national policy for hepatitis B birth dose vaccination in Philippines: lessons for improved delivery. Vaccine. 2011;29(5):941–5.
- MacDonald NE, Hesitancy SWGoV. Vaccine hesitancy: definition, scope and determinants. Vaccine. 2015;33(34):4161–4.

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Peter Francis Raguindin is a Vaccinologist and Epidemiologist who works on vaccine-preventable diseases and immunization system in the Philippines. He is a Senior Researcher at the Institute of Child Health and Human Development, National Institutes of Health – University of the Philippines, Manila, Philippines. He is also a Ph.D. student at the Institute of Social and Preventive Medicine - University of Bern, Bern, Switzerland.

Merrylle Morales-Dizon is a Pediatrician and a Senior Researcher at the Institute of Child Health and Human Development, National Institutes of Health – University of the Philippines, Manila, Philippines.

Josephine G. Aldaba is a Pediatrician and a Research Associate Professor at the Institute of Child Health and Human Development, National Institutes of Health – University of the Philippines, Manila, Philippines.

Lailani P. Mangulabnan is a Medical Officer at the Center for Health Development Region 3, Department of Health, San Fernando City, Pampanga, Philippines.

Renilyn P. Reyes is a Pediatrician and a Medical Officer at the Center for Health Development Region 6, Department of Health, Iloilo City, Iloilo, Philippines.

Batmunkh Nyambat is a Medical Officer for Vaccine Preventable Disease and Immunization at the World Health Organization Regional Office for the Western Pacific, Manila, Philippines.

Maria Joyce Ducusin is a Public Health Practitioner, who was the past Director of the Disease Prevention and Control Bureau, Department of Health, Manila, Philippines.

Anna Lena Lopez is a Pediatric Infectious Disease specialist and was the past Director of the Institute of Child Health and Human Development, National Institutes of Health, University of the Philippines, Manila, Philippines.



Affiliations

Peter Francis Raguindin^{1,2} · Merrylle Morales-Dizon¹ · Josephine Aldaba¹ · Lailani P. Mangulabnan³ · Renelyn P. Reyes⁴ · Batmunkh Nyambat⁵ · Maria Joyce Ducusin⁶ · Anna Lena Lopez¹

Merrylle Morales-Dizon mpmorales@up.edu.ph

Josephine Aldaba jgaldaba@up.edu.ph

Lailani P. Mangulabnan lpmangulabnan@gmail.com

Renelyn P. Reyes reyesrenilyn@yahoo.com

Batmunkh Nyambat batmunkhn@who.int

Maria Joyce Ducusin jducusin@gmail.com

Anna Lena Lopez allopez2@up.edu.ph

- Institute of Child Health and Human Development, National Institutes of Health University of the Philippines Manila, Manila, Philippines
- Present Address: Institute of Social and Preventive Medicine, University of Bern, Mittelstrasse 43, 3012 Bern, Switzerland
- Department of Health, Center for Health Development Region 3, San Fernando City, Pampanga, Philippines
- ⁴ Department of Health, Center for Health Development Region 6, Iloilo City, Iloilo, Philippines
- Vaccine Preventable Diseases and Immunization, World Health Organization Regional Office for the Western Pacific, Manila, Philippines
- Department of Health, Disease Prevention and Control Bureau, Manila, Philippines

