A retrospective study of acute pertussis in Hasan Sadikin Hospital—Indonesia

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**ABSTRACT**

**Objective:** To describe the representation of pertussis diagnosis in children. **Methods:** A retrospective observational study was performed on pediatric pertussis and pertussis—like syndrome registry for children <14 years of age documented from October 2008 to December 2014 in Hasan Sadikin Hospital, Indonesia. Demographic data, signs and symptoms at presentation, case definition (probable, confirmed), possible pertussis contact, pertussis vaccination status, results of *Bordetella pertussis* (*B. pertussis*) culture, complications, and outcome were recorded. **Results:** Sixty—one probable and two confirmed pertussis were documented. Male and female ratio was 1:1, mostly presented with shortness of breath, 24 (38%) subjects had posttussive vomiting, 10 (16%) had whooping—cough. Ten patients (16%) were reported to have adult possible pertussis contact. Only 2 infants had previous pertussis vaccination. All subjects presented in the second week of illness were all diagnosed as bronchopneumonia but two. The mean age was 6 months, ranging from 0—50 months. One subject required mechanical ventilation. *B. pertussis* culture was performed only in 35 (56%) subjects but positive only in two. There were no fatal cases, 55 (87%) including the subject who need mechanical ventilation had good outcome. **Conclusions:** Mostly patients were admitted on paroxysmal phase when no more active *B. pertussis* could be found from nasopharyngeal secret. A rigorous history taking particularly excessive cough, posttussive vomiting, and pertussis vaccination status need to be taken into account.

1. Introduction

Pertussis is a severe disease in children and remains as one of the 10 infant death causes worldwide. Reports show occurrence of about 10 million infected cases with almost 400 000 pertussis—related deaths annually[1]. Whooping cough usually manifests in wide spectrum of signs from asymptomatic presentation to a mild coughing illness in adolescents and adults who have partial immunity due to past vaccination or a full spectrum of symptoms in unimmunized patients. Physicians’ awareness and curiosity is needed to diagnose probable pertussis. In fact, confirmed pertussis need adequate laboratory culture examination. However, negative tests cannot be accepted as a definitive result to exclude pertussis. Therefore, it is necessary to apply improved laboratory diagnosis tests[2,3].

Diagnosis of pertussis is often delayed in children, especially adolescent due to the lack of classic symptoms (e.g. whooping cough doesn’t always present) and low
awareness of some physicians. While the patient has not been diagnosed and treated, he can be the source of transmission among the community for several weeks. In developing countries such as Indonesia, diagnosis of confirmed pertussis is difficult to be obtained. To diagnose confirmed pertussis, a specific Bordet Gengou or Regan Lowe culture is needed. Therefore, adequate laboratory examination is required. Other laboratory tests to confirm a pertussis infection such as PCR and ELISA are not always available, particularly in developing countries. The authors in this report discuss representation of pertussis diagnosis in children presented in our hospital[3].

2. Materials and methods

This retrospective observational study was performed in pertussis and pertussis–like syndrome registry in Child Health Department Hasan Sadikin Hospital from October 2008 to December 2014. All children less than 14 years who were suspected to have pertussis or pertussis–like syndrome were registered. We recorded their demographic data, signs and symptoms at presentation, case definition (probable, confirmed), possible pertussis contact, pertussis vaccination history, complications, and outcome. Complete physical examination were documented. They were tested for complete blood count, differential counting, aerobic microorganism blood culture and resistency, also nasopharyngeal Bordet Gengou culture. Complication and outcome were also recorded.

Suspected cases of pertussis had nasopharyngeal swab obtained for culture from the posterior nasopharynx. Nasopharyngeal swab specimens were obtained using a Dacron™ (not cotton) swab inserted slowly through the nostril to the posterior pharynx. Subjects’ head were immobilized, then nasopharyngeal swab were inserted gently into the nostril until the posterior nares was reached. The swab was left in place for up to 10 seconds. If resistance was encountered during insertion of the swab, we removed it and attempted insertion on the opposite nostril. Afterwards, we removed the swab slowly. The tube was labeled and immediately transported to the laboratory[4]. Bordet Gengou media was obtained from Bio Farma, a vaccine manufacturer nearby our hospital.

The pertussis registry were inputted in Microsoft Excel 2010. Characteristic data were presented in percentage.

3. Results

From October 2008 to December 2014, there were 61 probable and 2 confirmed pertussis cases in Child Health Department of Hasan Sadikin Hospital. Pertussis case occurrence was increasing in the last three years, with more than 10 diagnosed cases each year (Figure 1). The peak number of pertussis cases in our hospital was 17 cases in 2013.

The subjects’ mean age was 6 months (ranging from 1 to 50 months). Most subjects belonged to the age group of 0–5 months. Male to female ratio was 1:1. Sixty–one (96%) subjects were presented shortness of breath as chief complaint while the rest were presented with coughing. Six subjects (10%) had seizure. All of the subjects presented with coughing with durations ranging from 4 h to 1 month.

On admission, 28 (44%) subjects had been complaining about coughing for more than 1 week. Only ten (16%) subjects reported whooping cough. Twenty–four (38%) subjects reported posttussive vomiting while none of them had apneic symptoms. The average duration of symptoms before coming to the hospital was 11.8 days. Fifty–seven (90%) subjects had been treated with antibiotics, mostly amoxycillin by local health centres or general practitioners nearby. Only 2 infants had previous pertussis vaccination.

Forty–eight (76%) and 58 (92%) subjects were found to have leukocytosis and absolute lymphocytosis respectively. Bordet–Gengou agar cultures were performed in 35 (56%), with only 2 tests (3%) yielded positive results. Based on the classifications (probable and confirmed), only 2 (3%) subjects had confirmed pertussis while the rest were thought to have probable pertussis. Sixty–one (96%) subjects were initially diagnosed as having suspected bacterial pneumonia, while two were directly diagnosed as having pertussis. One subject had severe pneumonia and respiratory failure which required mechanical ventilation. Macrolide was given to all subjects, clarithromycin was given to 52 (83%) and the rest were given erithromycin. Most subjects showed good clinical responses, including the mechanical ventilated
subject. Although there was no fatal case, 10 (16%) subjects went home against medical advice. Previous diphtheria, tetanus and pertussis (DTP) vaccination were only received by 2 subjects (Table 1). History of respiratory tract infection in family which could be the possible contact was detected in 10 (16%) subjects.

Table 1
Characteristics of subjects in Child Health Department of Hasan Sadikin Hospital from October 2008 to December 2014.

<table>
<thead>
<tr>
<th>Characteristics of subjects (n=63)</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>26 (41%)</td>
</tr>
<tr>
<td>Female</td>
<td>37 (49%)</td>
</tr>
<tr>
<td>Age distribution</td>
<td></td>
</tr>
<tr>
<td>0–5 months</td>
<td>50 (79%)</td>
</tr>
<tr>
<td>6–11 months</td>
<td>6 (10%)</td>
</tr>
<tr>
<td>≥12 months</td>
<td>7 (11%)</td>
</tr>
<tr>
<td>Symptoms present</td>
<td></td>
</tr>
<tr>
<td>Cough</td>
<td>63 (100%)</td>
</tr>
<tr>
<td>Average length of cough, days</td>
<td>11.8</td>
</tr>
<tr>
<td>Fever</td>
<td>53 (84%)</td>
</tr>
<tr>
<td>Dyspnea</td>
<td>61 (96%)</td>
</tr>
<tr>
<td>Posttussive vomiting</td>
<td>24 (38%)</td>
</tr>
<tr>
<td>Laboratory findings</td>
<td></td>
</tr>
<tr>
<td>Leukocytosis</td>
<td>48 (76%)</td>
</tr>
<tr>
<td>Lymphocytosis</td>
<td>58 (92%)</td>
</tr>
<tr>
<td>Bordet Gengou culture</td>
<td>35 (56%)</td>
</tr>
<tr>
<td>Positive</td>
<td>2 (3%)</td>
</tr>
<tr>
<td>Negative</td>
<td>33 (97%)</td>
</tr>
<tr>
<td>Previous DTP immunization</td>
<td></td>
</tr>
<tr>
<td>Macrolide</td>
<td>63 (100%)</td>
</tr>
<tr>
<td>Clarithromycin</td>
<td>52 (83%)</td>
</tr>
<tr>
<td>Eritromycin</td>
<td>11 (17%)</td>
</tr>
<tr>
<td>Case definitions</td>
<td></td>
</tr>
<tr>
<td>Probable</td>
<td>61 (97%)</td>
</tr>
<tr>
<td>Confirmed</td>
<td>2 (3%)</td>
</tr>
<tr>
<td>On medical advice</td>
<td>53 (84%)</td>
</tr>
<tr>
<td>Hospital discharge</td>
<td></td>
</tr>
<tr>
<td>On medical advice</td>
<td>53 (84%)</td>
</tr>
<tr>
<td>Against medical advice</td>
<td>10 (16%)</td>
</tr>
</tbody>
</table>

4. Discussion

We documented 61 probable and 2 confirmed pertussis during the study period from Hasan Sadikin Hospital. There was increasing trend of pertussis numbers in our hospital in the last three years. The peak number was in 2013 with 17 cases and reduced in 2014. This may likely due to the start of our country’s new health insurance system which was held by Badan Penyelenggara Jaminan Sosial. Hasan Sadikin General Hospital is a tertiary hospital that is the center of national referral. Before coming to tertiary hospitals, all patients should have been treated in primary or secondary hospitals. They were only referred if the disease was considered complicated. As pertussis can be treated by antibiotics, the cases number might be increasing in primary or secondary hospital. The other reason may be undiagnosed pertussis. These might explain the lower pertussis incidence in the past year.

Most subjects (89%) were infants with the oldest age of 50 months. This is in contrary to the world’s latest epidemiology that reported the increase of pertussis prevalence in adolescents[8].

The Indonesian Ministry of Health reported 75.6% DTP vaccination coverage in 2013. Specifically in West Java, the coverage was 71.5%[9]. Only 2 subjects had previous pertussis vaccinations, which is very low as compared to the government’s report. This might be the reason why the peak incidence in our report was under 6 months of age. Anti-vaccination movement is also one of our problems nowadays. This may be one of many reasons why our subjects were not vaccinated. Apart from children’s vaccination coverage, the increasing number of adults’ pertussis may become the source of contact[4]. Until at present, Indonesia still doesn’t have Tdap vaccination program. This also explained the peak age incidence in our reports.

On the other hand, a resurgence of disease was also observed worldwide, with significant mortality in infants. Possible causes for this include the switch from whole-cell vaccines to less effective acellular vaccines, weaning immunity, pathogen adaptation, and increased use of more sensitive diagnostic tests in other countries[7]. Acellular vaccine formulations differ in the number of pertussis antigen component as compared to whole-cell vaccine. Some countries used vaccines which contain pertussis toxin (Ptx), pertactin (Prn), and filamentous hemagglutinin (Fha)[8]. Surveillance of recent Bordetella pertussis (B. pertussis) isolate in several country has detected prn mutation. Pertactin is a virulence factor of B. pertussis that promotes adhesion to tracheal epithelial cells. Whether Prn negative B. pertussis has greater or less virulence than positive strains still remains unclear. But the study in mice shows that loss of Prn does not seem to affect its lethality in mice[9]. In a retrospective study, no differences were found in the severity of symptoms or duration of hospitalization among infants infected with Prn-positive and Prn-negative strain. Regardless of Prn expression, vaccination reduced the severity of disease and the likelihood of being admitted to intensive care, which suggests that even an incomplete course of primary vaccination provides some protection against severe pertussis[10].

The peak of age was 0–5 months, ranging from 0–50 months. This data fits a review which states that this age group has the highest reporting rates and severity of disease, but is too young to be protected by the current vaccination schedule[11].
are also several journal reporting newborn with pertussis. The
youngest age was 18 days old. The baby had 2–day history of
cough without fever and worsening status. Both cases reported
that the mother had only mild cough[12,13]. One of the subjects
was 29 days old on admission.

We failed to detect the source of contact. The baby also had
the Bordet Gengou culture with negative result. The baby
was discharged with a good condition. We had no data for the
neonates because they are admitted to neonatology division.

All of the subjects were presented with cough, a main
symptoms of pertussis. Unfortunately, they all came in
paroxysmal phase (11.8 d in average). Only two of them was
diagnosed for having pertussis. The rest complained
and dyspnea, the reason they were diagnosed as
having bacterial pneumonia and were treated by intravenous
antibiotic. Twenty–four subjects reported postussive
vomiting. Rigorous anamnesis is very important in finding
the hallmark complaint of the disease.

Most of our subjects showed leukocytosis and absolute
lymphocytosis. This is in accordance with lymphocytosis–
promoting factor, a toxin protein of B. pertussis that induces
leukocytosis and lymphocytosis[14].

B. pertussis is a small Gram–negative cocccobacillus,
fastidious and is difficult to grow on media that is frequently
used in the laboratory to grow respiratory pathogens. B.
pertussis requires supplemental growth factors such as
Bordet–Gengou and Regan–Lowel[15].

Even though culturing B. pertussis is the gold standard, it
has low sensitivity and may be influenced by prior antibiotic
treatments, immunization status, duration of illness prior
to culture, specimen transport time, poor specimen quality,
and lack of laboratory expertise[16]. The Centers for Disease
Control and Prevention Pertussis Laboratory recommended
a Regan–Lowe medium because of its superiority over the
non–selective Bordet–Gengou medium[17]. A significant
advantage of RL is the 8–week shelf life at 4 °C, as compared
to 1 to 2 weeks for Bordet–Gengou. Regan–Lowe is also
superior at half strength for transport and enrichment[18].
Since our hospital could not do the culture, the subjects’
specimens had to be sent to the BioFarma Laboratory located
approximately 100 meters away. There were also no RL
medium, since all used Bordet–Gengou medium. Specimens
are collected after cough. Culture is best done in catarrhal
phase, which occurs during the first 2 weeks of symptoms
when bacteria are still present in the nasopharynx. After the
first 2 weeks, the sensitivity reduced and the risk of false
negative increased. Three weeks after commencement of
cough, culture sensitivity is only 1%–3%[19]. Our subjects
came to hospital after an average of 11.8 d of symptoms.
On the other hand, all subjects were borned by government
insurance which didn’t cover culture test. These reasons
explain not all the subjects’ specimen were sent for culture
and only 2 (3%) subjects yielded positive culture. The timing
of culture is generally in 2 weeks[15] but we usually obtained
the result in a week.

PCR is a rapid test and has excellent sensitivity and high
specificity which should be tested from nasopharyngeal
specimens taken at 0–3 weeks following cough onset, but may
provide accurate results for up to 4 weeks, 2 weeks longer
than culture. After the fourth week of cough, the amount of
bacterial deoxyribonucleic acid (DNA) diminishes rapidly,
which increases the risk of false negative results. Another
test to confirm pertussis diagnosis is serology. Optimal timing
for specimen collection is 2 to 8 weeks following cough
onset, when the antibody titers are at their highest. However,
serology may be performed on specimens collected up to 12
weeks following cough onset. Unfortunately, serology has
lower sensitivity and specificity as compared to PCR[20]. Both
PCR and serology are not available in our hospital and so was
in other laboratory in town.

Ampicillin or cephalosporin was given for bronchopneumonia
diagnosis while clarithromycin was given on the second week
following initial symptoms of most infants. Erythromycin was
given only to older infants to prevent hyperthropic pyloric
stenosis in young infant[21]. There were no fatal case and the
subject who required mechanical ventilation also had good
outcome, but some subjects went home against medical advice.

Factors that contribute to the difficulties in diagnosing
confirmed pertussis in our hospital were: (1) Most of our
patients were covered by government insurance, which do
not include culture for B. pertussis; (2) Most subjects came at
the paroxysmal phase when no more active B. pertussis could
be found from nasopharyngeal secret; (3) PCR and ELISA
were not provided in our laboratory.

There were several limitations of this study. We had no
data regarding immunization statuses of the parents and
clinical and laboratory examination. Therefore, we could
not find out whether they were the household contacts.
Case findings at primary and secondary health care and
influencing factors analysis are really encouraged in order to
describe the real magnitude of pertussis.

Mostly patients were admitted during paroxysmal phase
when very limited number of *B. pertussis* could be identified from nasopharyngeal secret. A rigorous history taking, particularly excessive coughing, posttussive vomiting, and pertussis vaccination status need to be taken into account.

**Conflict of interest statement**

The authors report no conflict of interest.

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**References**


