Evaluation of immunisation strategies for pertussis vaccines in Jinan, China – an interrupted time-series study

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Abstract

Studies in countries with high immunisation coverage suggest that the re-emergence of pertussis may be caused by a decreased duration of protection resulting from the replacement of whole-cell pertussis vaccine (WPV) with the acellular pertussis vaccine (APV). In China, WPV was introduced in 1978. The pertussis vaccination schedule advanced from an all-WPV schedule (1978–2007), to a mixed WPV/APV schedule (2008–2009), then to an all-APV schedule (2010–2016). Increases in the incidence of pertussis have been reported in recent years in Jinan and other cities in China. However, there have been few Chinese-population-based studies focused on the impact of schedule changes. We obtained annual pertussis incidences from 1956 to 2016 from the Jinan Notifiable Conditions Database. We used interrupted time series and segmented regression analyses to assess changes in pertussis incidence at the beginning of each year, and average annual changes during the intervention. Pertussis incidence decreased by 1.11 cases per 100 000 population (P = 0.743) immediately following WPV introduction in 1978 and declined significantly by 1.21 cases per 100 000 population per year (P < 0.0001) between 1978 and 2001. Immediately after APV replaced the fourth dose of WPV in 2008, the second and third doses in 2009, then replaced all four doses in 2010, pertussis incidence declined by 1.98, 1.98 and 1.08 cases per 100 000 population, respectively. However, the results were not statistically significant. There were significant increasing trends in pertussis incidence after APV replacements: 1.63, 1.77 and 1.78 cases/year in 2008–2016, 2009–2016 and 2010–2016, respectively. Our study shows that the impact of an all-WPV schedule may be less than the impacts of the sequential WPV/APV schedules. The short-term impact of APV was better than that of WPV; however, the duration of APV-induced protection was not ideal. The impact and duration of protective immunity resulting from APVs produced in China need further evaluation. Further research on the effectiveness of pertussis vaccination programme in Jinan, China is also necessary.

Introduction

Pertussis is a worldwide, cyclic acute respiratory infection caused by Bordetella pertussis. Prior to widespread coverage of the pertussis vaccine (PV), pertussis was the primary cause of death among infants and young children [1]. The PV resulted in a marked decrease in the morbidity and mortality rates since the 1950s [1]. However, despite widespread implementation of these different immunisation programmes and associated levels of uptake, pertussis continues to persist. Since the 1980s, there has been increased incidence and shifts in age-specific incidence towards a higher representation of adolescents and adults have been reported in many developed countries with high vaccine coverage, which has been labelled so-called ‘re-emergence of pertussis’. Available evidence indicates that there is a more rapid waning of immunity, and possibly a reduced impact on transmission, with acellular pertussis vaccine (APV) relative to whole-cell pertussis vaccine (WPV) [1–3].

To further control pertussis, the International Consensus Group on Pertussis Immunization [4] and the Global Pertussis Initiative [5] have both recommended a universal adolescent booster vaccination combined with targeted immunisation of those adults most likely to have contact with babies, including parents, close family members, healthcare workers and day-care workers; universal immunisation of young adults and routine use of tetanus–diphtheria–pertussis vaccine (Tdap) rather than tetanus toxoid and diphtheria vaccine (Td) for all adult booster immunisations. In Canada, Germany, Belgium and the United States, two booster shots of APV products are recommended for 4–7-year-old children and 11–17-year-old adolescents after four doses of routine vaccination [6]. In Australia, a cocoon programme offering parents of new babies a government-funded dose of a pertussis-containing vaccine was implemented [7]. In the United States, the United Kingdom, Switzerland and Australia, a maternal vaccination programme was offered to prevent cases in young infants [8].
In China, pertussis vaccination prepared from WPV was available to children as part of the publicly funded National Immunization Programme from 1978 through 2007, protecting infants who were 2 months of age or older. Due to safety and effectiveness, APV replaced the whole-cell product in 2008. Vaccination against pertussis was recommended at 3, 4 and 5 months, and between 18 and 24 months of age. The vaccination coverage survey conducted in 2008 showed that in children aged 1–2 in China, 99.10% had an immunisation certificate, and 99.38% had an immunisation card, with an overall vaccination coverage of 99.42% [9].

The incidence of pertussis in China decreased from between 100 and 200 cases per 100,000 population during the pre-vaccination era of the 1960s and 1970s [10] to a historic low of less than one case per 100,000 population from 2004 to 2016 [11–13]. In China, pertussis cases were usually identified via passive hospital reports. Although two types of diagnostic criteria [14, 15] were used in 1995–2006 and 2007–2016, respectively, there were only minor changes to surveillance methods in China during these time periods. Most cases were clinically diagnosed and laboratory methods were not routinely used in hospitals [16].

As the capital city of Shandong province in the east of China, Jinan achieved the goal of immunisation coverage among 85% of targeted children at county and township levels in 1991 and 1996, respectively. From then on, PV coverage in Jinan has consistently been greater than 95% (CDC, unpublished data). According to the national immunisation schedule, WPV was introduced into the Expanded Programme on Immunisation (EPI) as a government-purchased vaccine (category 1 vaccine) from 1978 to 2009 in Jinan. Category 1 vaccines refer to free vaccines that are mandatory for children in China to be vaccinated in accordance with the government’s EPI. Between 2002 and 2007, APV was introduced as a category 2 vaccine that refers to vaccines that are not for routine use, are optional for residents and must be privately bought in China. In 2008, APV became a category 1 vaccine, making it free for the fourth dose, and WPV was still in use for prior doses one to three. In 2009, WPV was free for the first dose and APV for the following two to four doses. From 2010 to the present time, only APV is available. Before the PVs were introduced from 1956 through to 1977, the mean reported incidence of pertussis in Jinan was 67.61 cases per 100,000 population, with a peak value of 303.93 in 1963. During the vaccination era from 1978 to 2016, the incidence decreased from 14.84 cases per 100,000 population to a historic low of 0.02 per 100,000 in 2003, increased to 3.08 in 2014, then continued to increase to 4.74 in 2015 and 6.85 in 2016.

The reasons for the re-emergence of pertussis in Jinan from 2014 through 2016 are not well understood. In previous studies, protection effectiveness of WPV and APV was mostly calculated using case-control studies and field trials of mass vaccination [3, 17–22]. In China, there were only case-control studies of immunogenicity and immune persistence of APV with small sample sizes. Therefore, further research on the pertussis vaccination programme in Jinan, China is necessary. In this study, we used a long-term macroscopic evaluation to examine the impact of PV schedule changes. Interrupted time series (ITS) design and segmented regression analysis were utilised to analyse PV immunisation strategies based on the trends of pertussis incidence rates from 1956 through to 2016 in Jinan.

Methods

Study design

As a large-scale intervention, it is difficult to evaluate the effect of a mass vaccination intervention by a randomised controlled trial on the population level without randomisation or any control. We conducted a longitudinal study of the reported pertussis incident between 1956 and 2016 with a single group of ITS design for evaluating the impacts before and after mass vaccination interventions.

Data source

This study was conducted using the data of annually reported pertussis incidence in Jinan. The data of reported cases and permanent population in 1956–2003 was collected from the Annual Report on the Incidence of Infectious Diseases in Jinan City. Data from 2004–2016 came from the Infectious Disease Reporting Information Management System, one subsystem of China’s Information System for Disease Control and Prevention.

Case definition

The cases were all hospital reported patients living in Jinan City for more than 3 months. In China, two types of diagnostic criteria were used in 1995–2006 and 2007–2016, respectively. In 1995 [14], the clinical case definition required one of the symptoms to be either a cough for at least 14 days, a paroxysmal spasmodic cough, a cough with vomiting, unknown paroxysmal asphyxia in infants or on the basis of an epidemiologic link. This confirmation could be based on laboratory results, for example, a positive culture for B. pertussis or a significant fourfold increase in a specific serum antibody. In 2007 [15], a suspected case was confirmed with an acute cough lasting more than 14 days, paroxysms of coughing, inspiratory ‘whoop’, posttussive vomiting or direct contact (epidemiology linkage). A clinical case was defined by a suspected case with a significantly increasing calculation of leucocyte and lymphocyte in peripheral blood. The laboratory-confirmed case was confirmed as a clinical case with isolation of B. pertussis from phlegm or nasopharyngeal secretion, or the serum specific antibody of the convalescence period was four times higher than the acute phase.

Outcome variable

Y, was the main outcome variable that corresponds to the annual incidence rates of hospital reported pertussis, including suspected, clinical and laboratory-confirmed cases. Annual pertussis incidences were calculated at each time point by the number of cases per year obtained from the resident population, divided by the resident population in the year, then multiplied by 100,000 inhabitants.

Intervention definitions

According to the immunisation schedules in Jinan, the history of intervention can be subdivided into seven stages from 1956 to 2016. Stage A was the pre-vaccination period from 1956 to 1977. Stage B was the timeframe of which the WPV vaccination programme was introduced. Annual average coverage during this period was 67.61%. Stage C was a WPV vaccination period between 1991 and 2001, with an average coverage of more than 90%.
90%. Stage D, between 2002 and 2007, was the WPV vaccination period, with the introduction of APV as a category 2 vaccine. In stage E, APV was introduced as a category 1 vaccine for the fourth dose in 2008. In stage F, APV replaced WPV for doses 2 through to 4 as the category 1 vaccine in 2009. In stage G, APV replaced all four doses of WPV from 2010 to 2016 (Table 1).

Vaccination interventions were identified into five statuses according to the shifts of immunisation strategy including (1) Intervention 1: stage A (1956–1977) was the pre-intervention area and stages B and C (1978–2001) were considered as the WPV intervention stage. We regarded the year 1978 as the PV intervention time point. (2) Intervention 2: stage B (1991–2001) was considered as the WPV-intervention period with coverage of more than 90%. Stage C (2002–2007) was set as the APV (self-funded vaccine and low coverage) vaccination intervention stage. Year 2002 was regarded as the beginning of APV vaccination intervention. (3) Intervention 3: stages C and D were considered as the pre-intervention period with WPV vaccination. Stages E–G were reckoned as the APV vaccination intervention era. Year 2008 was regarded as the beginning of APV vaccination intervention for the replacement of APV for the fourth dose of WPV. (4) Intervention 4: stages C–E (1991–2008) were considered the pre-intervention period of the WPV vaccination and stages F and G (2009–2016) as the APV vaccination intervention stage. Year 2009 was regarded as the beginning of APV vaccination intervention for the replacement of APV for the second–fourth dose of WPV. (5) Intervention 5: stages C–F (1991–2009) were considered as the pre-intervention period of the WPV vaccination and stage G as the APV vaccination intervention stage. Year 2010 was regarded as the beginning of all four doses of APV vaccination intervention (Table 1).

**Statistical analysis**

The segmented regression model is most commonly used to assess ITS data. It is fitted through to a least squares regression line to each segment of the independent variable and time, thus assuming a linear relationship between time and the outcome within each segment [23, 24]. We used segmented regression analysis to assess the significance of changes in levels and trends of reported pertussis incidence before and after the vaccination interventions, including implementation of PV and alternative strategies. The level is the value of the series at the beginning of a given time interval (i.e. the y-intercept for the first segment and the value immediately following each change point at which successive segments join). A change in level constitutes an abrupt intervention effect. The trend is the rate of change of a measure (in other words, the slope) during a segment. A change in trend is defined by an increase or decrease in the slope of the segment after the intervention as compared with the segment preceding the intervention. The regression model was set as follows:

\[ Y_t = \beta_0 + \beta_1 \times \text{Year} + \beta_2 \times \text{ Intervention} + \beta_3 \times \text{Postslope} + \epsilon_t \]

where \( Y_t \) is the reported pertussis rate per year. Year was a continuous variable indicating time in years from the beginning of the observation period at time (1). We used log 10-transformation first, then used a moving average to pre-process the raw incident data, which can be seen in Figure 1. This was completed to remove the cycles and better capture the trends. Then we analysed using interrupted time-series analysis [25]. Intervention was an indicator variable for time (t) occurring either before (intervention = 0) or after (intervention = 1) the mass vaccination. Postslope was a continuous variable counting the number of years at time (t) after the intervention, which was equal to Year × Intervention. \( \beta_0 \) estimates the baseline level of the outcome at time zero. \( \beta_1 \) estimates the overall linear trend of yearly change in the pertussis rate. \( \beta_2 \) estimates the change in level in the rate immediately after the intervention from the end of the preceding stage. \( \beta_3 \) estimates the long-term trend change in the outcome after the intervention, compared with the trend before the intervention. \( \epsilon_t \) is the error term at time (t), representing the random variability not explained by the model. The Durbin–Watson (DW) method was applied to test the stability of time series data.

| Stage | Policy period | Average incidence (per 100,000) | Incidence range (per 100,000) | P-value* | Average coverage (%) | Coverage range (%) | Immunisation schedule |
|-------|---------------|-------------------------------|-------------------------------|----------|-----------------------|--------------------|----------------------|
| A     | 1956–1977     | 67.61                         | (2.10–303.93)                 | <0.0001  | Unknown               | Unknown            | Pre-vaccination period |
| B     | 1978–1990     | 4.80                          | (0.47–14.84)                  | <0.0001  | 75.71                 | (65.67–87.54)      | 4 doses of WPV vaccination |
| C     | 1991–2001     | 0.27                          | (0.05–0.62)                   | <0.0001  | 90.52                 | (86.07–94.07)      | WPV as category 1 vaccine and APV as category 2 vaccine |
| D     | 2002–2007     | 0.11                          | (0.02–0.21)                   | 0.068    | 96.13                 | (94.77–97.32)      | 4 doses of WPV vaccination |
| E     | 2008          | 0.18                          | –                            | <0.0001  | 96.71                 | –                  | 1–3 doses of WPV and the 4th dose of APV |
| F     | 2009          | 0.03                          | –                            | <0.0001  | 98.45                 | –                  | 1 dose of WPV and 2–4 doses of APV |
| G     | 2010–2016     | 2.58                          | (0.23–6.85)                   | 99.53    | (98.34–99.83)         | 4 doses of APV vaccination |

*P value of \( \chi^2 \) test: the average of incidence compared with the next period.
To describe how the immunisation strategies affect the all-age pertussis incidence, we analysed the incidences of birth cohorts from 2005 to 2016 (Table 2). All statistical analyses were conducted using SAS statistical software (ver 9.3). The statistical significance level was set at 0.05.

**Results**

In the pre-vaccination era, the reported incidence of pertussis ranged between 2.10 and 303.93 cases per 100 000 population during 1956–1977, with an average incidence of 67.61 cases per 100 000, which reduced to an average value of 4.80 per 100 000 population with a range between 0.47 and 14.84 during 1978–1990, reduced to 0.27 (0.05–0.62) in 1991–2001 and 0.11 (0.02–0.21) in 2002–2007. In the period of the shift from WPV to APV, the incidence was 0.18 in 2008 and 0.03 in 2009, respectively. In the APV vaccination intervention stage (2010–2016), the mean value of incidence raised to 2.58 (0.23–6.85). There were statistically significant differences between the average incidence of one policy period and the subsequent period, except for the difference

**Fig. 1.** Pertussis reported incidence rates in Jinan, 1956–2016. Blue line depicts the trend in log-transformed per capita pertussis incidences in Jinan, China from 1956 to 2016. Orange line depicts smoothing spline in backward moving average with five periods. Seven colour shaded regions in the figure present stages A–G with different immunisation schedules consistent with Table 1.

**Table 2.** The pertussis incidences of birth cohorts from 2005 to 2016 (per 100 000)

| Birth cohort | 2005 | 2006 | 2007 | 2008 | 2009 | 2010 | 2011 | 2012 | 2013 | 2014 | 2015 | 2016 |
|--------------|------|------|------|------|------|------|------|------|------|------|------|------|
| 2005         | 2.80 | 1.42 | 0.00 | 0.00 | 1.42 | 0.00 | 0.00 | 0.00 | 0.00 | 5.77 | 5.13 | 0.00 |
| 2006         | 2.80 | 0.00 | 0.00 | 0.00 | 1.42 | 0.00 | 1.61 | 0.00 | 5.87 | 1.91 | 24.57|
| 2007         | 4.18 | 2.82 | 0.00 | 0.00 | 0.00 | 1.60 | 0.00 | 10.93| 0.00 | 28.81|      |
| 2008         | 4.15 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 3.20 | 13.07| 0.00 | 10.56|      |
| 2009         | 1.38 | 4.19 | 0.00 | 3.12 | 1.56 | 25.22| 9.75 | 35.51|      |
| 2010         | 35.66| 7.46 | 0.00 | 1.56 | 18.60| 4.70 | 37.54|      |
| 2011         | 67.46| 0.00 | 7.76 | 22.97| 10.79| 51.98|      |
| 2012         |      |      |      |      |      |      |      |      |      |      |      |      |
| 2013         |      |      |      |      |      |      |      |      |      |      |      |      |
| 2014         |      |      |      |      |      |      |      |      |      |      |      |      |
| 2015         |      |      |      |      |      |      |      |      |      |      |      |      |
| 2016         |      |      |      |      |      |      |      |      |      |      |      |      |
between 2002–2007 and 2008. Table 1 shows the average coverage rose from 75.71% of three doses of PV during 1978–1990, to 90.52% during 1991–2001 and reached 99.53% in 2010–2016.

Figure 2 shows the distribution of annual pertussis cases by five different age groups (under 1 year, 1–2, 3–6, 7–14 and 15–59 years). From 2005 to 2016, the age distribution of cases remained relatively stable with between 83.33% and 100% of all cases attributed to infants younger than 6-year-old, especially the cases under 1 year. In the APV vaccination intervention stage (2010–2016), the percentage of cases under 1 year dropped from 86.67% in 2010 to 28.83% in 2016, with increasing proportions of cases in those aged 1–6 years from 13.33% in 2010 to 56.24% in 2016. Cases among those aged 7–14 and 15–59 years had been reported since 2014, with the cumulative percentage of cases in both age groups ranging from 8.74% to 14.93% (Fig. 2).

Intervention 1: WPV vaccination as the intervention in 1978–2001

As presented in Table 3, the segmented regression analysis results indicate that before the vaccination intervention era, the trend of reported incidence of pertussis declined by 1.10 cases per 100 000 population every year ($P = 0.049$). The significant declining trends existed in every model by approximately one case per 100 000 population. When WPV was introduced in 1978, there was a slight decline in the incidence of 1.11 cases per 100 000 population ($P = 0.0743$) immediately. This statistically significant trend continued to 2001 when incidence decreased by 1.21 cases per 100 000 population per year ($P < 0.0001$) (Table 3 and Fig. 3a).

Intervention 2: the introduction of APV (as the category 2 vaccine) as the intervention in 2002–2007

In 2002–2007, APV was introduced as the category 2 vaccine. In 2002, there was a slight decrease in the level of incidence of 1.06 cases per 100 000 population, and an increase in trend in the incidence of 1.06 cases per 100 000 population from 2002 through 2007. The result indicated that there was no significant change in level ($P = 0.703$) and trend ($P = 0.267$) after the intervention (Table 3 and Fig. 3b).

Intervention 3: APV vaccination (WPV for 1–3 doses and APV for fourth dose in 2008) as the intervention in 2008–2016

In 2008, APV was introduced as the category 1 vaccine for the fourth dose. There was an immediate decline in the incidence of 1.98 cases per 100 000 population and there was no significance ($P = 0.774$). However, there was a significant increasing trend from 2008 through 2016 of 1.63 cases per 100 000 population per year ($P < 0.0001$) (Table 3 and Fig. 3c).

Intervention 4: APV vaccination (WPV for 1 dose and APV for doses 2–4 in 2009) as the intervention in 2009–2016

In 2009, when APV replaced WPV for the doses 2–4, the incidence declined immediately by 1.98 cases per 100 000 population ($P = 0.773$). However, the trend increased by 1.77 cases per 100 000 population per year and was significant ($P < 0.0001$) from 2009 to 2016 (Table 3 and Fig. 3d).

Intervention 5: APV vaccination as the intervention in 2010–2016

When APV replaced WPV for doses 1–4 in 2010, there was an immediate decrease in the incidence of 1.08 cases per 100 000 population ($P = 0.733$). There was a significant increasing trend in the incidence by 1.78 cases per year from 2010 through 2016 ($P < 0.0001$) (Table 3 and Fig. 3e).

The incidence in the birth year of each cohort increased from 2.80 per 100 000 in the 2005 cohort, and to 182.31 per 100 000 in the 2016 cohort. After the last receipt of all-WPV schedules, the incidence was lower than 3 cases per 100 000 population. This lasted for 8 years in the 2005 cohort, 7 years in the 2006 cohort and 6 years in the 2007 cohort. However, in the 2008 cohort the incidence began to increase in 2013, 4 years after the last receipt of APV. In the 2009–2011 cohort, the lasting years after the fourth dose and before the increase of incidence was only 1–2 years. Two years after the all-APV schedule was recommended in 2012, the incidences further increased for all cohorts in 2014 and the periods of protection effect were inapparent. In 2016, yearly incidences for every cohort reached their peak (Table 2).
high level of protection against pertussis \[1\].

or APV, regardless of vaccine type or manufacturer, can provide a scientific pieces of evidence demonstrate that a primary series of WPV could effectively reduce the incidence of pertussis. Numerous scientific pieces of evidence demonstrate that a primary series of WPV mass vaccination from 1978 to 2001, respectively. It indicated that WPV mass vaccination was significantly lower in children with basic immunisation and immune-strengthening using APV than in WPV \[1\].

was increased compliance to the vaccination schedule, promoted by the perceived safety of APV, as this would guarantee vaccine efficacy \[6\]. Related studies found that the incidence of side effect increased at the rate of 1.63 per 100 000 population with the intervention from 2001; after: 2002–2007

2016, respectively. It is chosen as the strongest quasi-experimental approach for evaluating the longitudinal effects of interventions. Therefore, outcomes before and after implementation of the intervention were collected and compared, while accounting for potential confounders and potential secular data trends \[26, 27\].

In the model for the intervention of WPV mass vaccination, there were level and trend changes that decreased at the rate of 1.11 and 1.21 per 100 000 population with the intervention from 1978 to 2001, respectively. It indicated that WPV mass vaccination could effectively reduce the incidence of pertussis. Numerous scientific pieces of evidence demonstrate that a primary series of WPV or APV, regardless of vaccine type or manufacturer, can provide a high level of protection against pertussis \[1\].

Our study found that a combination of 1–3 doses WPV and a fourth dose APV in 2008, and a combination of 1 dose WPV and 2–4 doses APV in 2009 had the highest impacts, with an immediate level decrease of 1.98 cases per 100 000 population in 2008 and 2009, respectively. The results indicate that the impact of a mixed schedule including both WPV and APV was better than four doses of only WPV. This can be interpreted as a result of increased compliance to the vaccination schedule, promoted by the perceived safety of APV, as this would guarantee vaccine efficacy \[6\]. Related studies found that the incidence of side effect was significantly lower in children with basic immunisation and immune-strengthening using APV than in WPV \[1\].

Despite decreasing level changes in the annual rate immediately after the APV intervention, compared to the WPV vaccination era, changes in trend were increasing at the rate of 1.63 cases per 100 000 population in 2008–2016, 1.77 cases per 100 000 population in 2009–2016 and 1.78 cases per 100 000 population in 2010–2016, respectively. The changes in trend may be associated with the WPV-induced immunity persistence that is longer than APV’s \[28\]. Evidence from a systemic meta-analysis suggests that the average duration of protective immunity to pertussis after the fifth dose of APV is 3–4 years \[29\]. However, a mathematical model analysis has shown a longer duration of DTaP immunity in the United States than the meta-analysis. The result demonstrates that APV confers imperfect, but has long-lived protection \[30\]. There is also ample evidence that APV confers long-term protection \[31–34\].

Although globally there are a wide variety of APV vaccination schedules that are recommended, for every additional year after the last dose of APV, the odds of pertussis increase by 1.33 times \[29\]. A multi-centre clinical trial showed that the efficacies of most APV products in America were significantly different, which suggested that the source, method of manufacture and included antigens of the vaccine should be considered in the comparative study \[35\]. In China, the now available APV consists predominantly of FHA and inactivated PT and are known as Takeda-type vaccines \[36\]. A study of the serological effects of two types of PV developed by Chinese producers showed that higher levels of anti-PT antibody and anti-FHA antibody were detected in sera from children boosted with three doses APV than that boosted with three doses WPV in 1 month and 6 months after vaccination \[37\].

In the all-WPV era, the imperfect, but nevertheless effective vaccines conferred a slow waning protection and strong herd immunity. In the 2005–2007 cohort, a shorter phase of low

**Table 3.** Estimated level and trend changes of reported pertussis incidence before and after PV vaccination inventions

| Intervention model                     | Coefficient  | S.E.  | P-value | DW  | R²  |
|----------------------------------------|--------------|-------|---------|-----|-----|
| (1) Before: 1960–1977; after: 1978–2001| Intercept β₀ | 77.65 | 1.80    | <0.0001 | 0.760 | 0.703 |
|                                        | Baseline trend β₁ | −1.10 | 1.05    | 0.049 |
|                                        | Level change after intervention β₂ | −1.11 | 1.38    | 0.743 |
|                                        | Trend change after intervention β₃ | −1.21 | 1.03    | <0.0001 |
| (2) Before: 1991–2001; after: 2002–2007| Intercept β₀ | 1.50  | 1.15    | 0.012 | 1.465 | 0.869 |
|                                        | Baseline trend β₁ | −1.16 | 1.02    | <0.0001 |
|                                        | Level change after intervention β₂ | −1.06 | 1.16    | 0.703 |
|                                        | Trend change after intervention β₃ | 1.06  | 1.05    | 0.267 |
| (3) Before: 1991–2007; after: 2008–2016| Intercept β₀ | 4.18  | 1.49    | 0.002 | 1.714 | 0.845 |
|                                        | Baseline trend β₁ | −1.15 | 1.02    | <0.0001 |
|                                        | Level change after intervention β₂ | −1.98 | 1.26    | 0.774 |
|                                        | Trend change after intervention β₃ | 1.63  | 1.05    | <0.0001 |
| (4) Before: 1991–2008; after: 2009–2016| Intercept β₀ | 2.33  | 1.5     | 0.05  | 1.741 | 0.833 |
|                                        | Baseline trend β₁ | −1.12 | 1.02    | <0.0001 |
|                                        | Level change after intervention β₂ | −1.98 | 1.25    | 0.773 |
|                                        | Trend change after intervention β₃ | 1.77  | 1.05    | <0.0001 |
| (5) Before: 1991–2009; after: 2010–2016| Intercept β₀ | 2.47  | 1.44    | 0.021 | 1.822 | 0.849 |
|                                        | Baseline trend β₁ | −1.12 | 1.02    | <0.0001 |
|                                        | Level change after intervention β₂ | −1.08 | 1.25    | 0.733 |
|                                        | Trend change after intervention β₃ | 1.78  | 1.06    | <0.0001 |
incidence after the last dose of WPV can be observed. This could be due to the epidemic cycle. As the cohort population is growing older, they may act as a reservoir of transmission to young children. Substantial pertussis circulation among adults and adolescents would inevitably lead to the increased incidence among infants, especially those too young to get immunised [34, 38]. When APV sequentially replaced WPV since 2008, the lasting low incidence after APV vaccination began growing shorter in the 2008–2016 cohort. Since 2014, incidences in all cohort had further increased, which may lead to the speculation that APV usage in China provided short-lived immunity. The high transmissibility of pertussis [2], the ‘end-of-honeymoon’ effect of WPV vaccination [6], persisted circulation [39] and the relatively high risk of disease in groups with high contact rates, such as schoolchildren [31] may also be cause for the increasing incidences of pertussis in Jinan city in recent years. The association between pertussis vaccination policy and incidences is intricate, making their interpretation difficult. However, the individual variability of primary vaccine failure and vaccine duration failure of APV should be considered as well [30].

Although surveillance data may underestimate disease incidence, biases would be expected to apply equally to local reporting and vaccination patterns over time. At the same time, this study did not control potential variables including social and economic factors, migration [40], vaccine production and only considered factors such as the implementation of different vaccines and vaccination programmes. Multi-factor analysis should further be researched.

The results of this study show that the impact of WPV may be less than that of a combination of WPV and APV, and the short-term influence of APV is better than that of WPV; however, the duration of APV-induced immunity is not ideal. Analogous to the developed countries with high vaccine coverage, the incidence of pertussis has increased in Jinan. As long as currently available
APVs are in use, it is likely that the ‘new normal’ will be higher disease incidence throughout pertussis cycles [39]. Furthermore, some inventions in other countries that had a similar rise in pertussis incidence had a reference value. In the UK, a maternal vaccination programme was offered to prevent cases in young infants who were not yet eligible for the vaccine [8]. The International Consensus Group on Pertussis Immunization [4] and the Global Pertussis Initiative [5] both have recommended universal adolescent booster vaccination combined with targeted immunisation of those adults most likely to have contact with babies, including parents close family members, healthcare workers and day-care workers; universal immunisation of young adults and routine use of Tdap rather than Td for all adult booster immunisations. However, despite the limitations of current PV, timely and complete immunisation with available PV is still the best practice against pertussis. Additional evaluation of vaccines and development of improved PVs and vaccination schedules in Jinan, China is necessary.

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References
1. World Health Organization (2015) Pertussis vaccines: WHO position paper. Vaccine 90, 433–458.
2. Wood N et al. (2008) Pertussis: review of epidemiology, diagnosis, management and prevention. Paediatric Respiratory Reviews 9, 201–211; quiz 211–202.
3. Tartof SY et al. (2013) Waning immunity to pertussis following 5 doses of DTaP. Pediatrics 131, e1047–e1052.
4. Campinsmartí M et al. (2001) Recommendations are needed for adolescent and adult pertussis immunisation: rationale and strategies for consid-eration. Vaccine 20, 641–646.
5. Forsyth KD et al. (2004) New pertussis vaccination strategies beyond infancy: recommendations by the global pertussis initiative. Clinical Infectious Diseases 39, 1802–1809.
6. Plotkin SA et al. (2011) Vaccines. Beijing: People’s Medical Publishing House.
7. Rowe SL et al. (2018) Effectiveness of parental cooing as a vaccination strategy to prevent pertussis infection in infants: a case-control study. Vaccine 36(15), 2012–2019.
8. Aretà ML et al. (2017) Antenatal vaccination to decrease pertussis in infants: safety, effectiveness, timing, and implementation. Journal of Maternal-Fetal and Neonatal Medicine 32, 1–6.
9. Cao L et al. (2012) National immunization coverage survey in China after integrated more vaccines into EPI since 2008. Chinese Journal of Vaccines and Immunization 18, 419–424, 478.
10. Li YF et al. (2017) Research progress on the effectiveness and persistence of pertussis vaccine induced immunity. Chinese Journal of Vaccines and Immunization 23, 230–233.
11. Ning GJ et al. (2018) Epidemiology of pertussis in China, 2011–2017. Chinese Journal of Vaccines and Immunization 24, 264–267, 273.
12. Wang HB et al. (2012) Epidemiological analysis on pertussis in China during 2006–2010. Chinese Journal of Vaccines and Immunization 3, 207–210.
13. Yin DP et al. (2007) Epidemiological analysis on pertussis in China during 2004–2006. Chinese Journal of Vaccines and Immunization 13, 245–247.
14. The Ministry of Health of People’s Republic of China. 1995. Diagnostic criteria and management principles of pertussis (in Chinese). Available at http://www.nhc.gov.cn/wjw/s9491/201212/34026.shtml.
15. The Ministry of Health of People’s Republic of China. 2007. Diagnostic Criteria for Pertussis (in Chinese). Available at http://www.nhc.gov.cn/wjw/s9491/201410/52040bc163db4eebaceae56ec28b335866.shtml.
16. Huang H et al. (2014) Epidemiological features of pertussis resurgence based on community populations with high vaccination coverage in China. Epidemiology and Infection 143, 1–7.
17. Cherry JD (2012) Waning protection after fifth dose of acellular pertussis vaccine in children. New England Journal of Medicine 367, 1012–1019.
18. Missagades LK et al. (2012) Association of childhood pertussis with receipt of 5 doses of pertussis vaccine by time since last vaccine dose, California, 2010. JAMA 308, 2126–2132.
19. Liko J et al. (2013) Priming with whole-cell versus acellular pertussis vaccine. New England Journal of Medicine 368, 581–582.
20. Gustafsson L et al. (1996) A controlled trial of a two-component acellular, a five-component acellular, and a whole-cell pertussis vaccine. New England Journal of Medicine 334, 349–355.
21. Schmitt HJ et al. (1996) Efficacy of acellular pertussis vaccine in early childhood after household exposure. JAMA 275, 37–41.
22. Stehr K et al. (1998) A comparative efficacy trial in Germany in infants who received either the Lederle/Takeda acellular pertussis component DTP (DTaP) vaccine, the Lederle whole-cell component DTP vaccine, or DT vaccine. Pediatrics 101, 1–11.
23. van Seben R et al. (2016) Implementation of a transfer intervention procedure (TIP) to improve handovers from hospital to home: interrupted time series analysis. BMC Health Services Research 16, 479.
24. Taljaard M et al. (2014) The use of segmented regression in analysing interrupted time series studies: an example in pre-hospital ambulance care. Implementation Science 9, 77.
25. Rohani P et al. (2011) The decline and resurgence of pertussis in the US. Epidemics 3, 183–188.
26. Wagner AK et al. (2002) Segmented regression analysis of interrupted time series studies in medication use research. Journal of Clinical Pharmacy and Therapeutics 27(4) 299–309.
27. Bernal JL et al. (2017) Interrupted time series regression for the evaluation of public health interventions: a tutorial. International Journal of Epidemiology 46, 348–355.
28. Snits K et al. (2013) Different T cell memory in preadolescents after whole-cell or acellular pertussis vaccination. Vaccine 32, 111.
29. McGirr A et al. (2015) Duration of pertussis immunity after DTaP immunization: a meta-analysis. Pediatrics 135, 331–343.
30. Domenech de Celles M et al. (2019) Duration of immunity and effectiveness of diphtheria-tetanus-acellular pertussis vaccines in children. JAMA Pediatrics 173, 588–594.
31. Domenech de Celles M et al. (2018) The impact of past vaccination coverage and immunity on pertussis resurgence. Science Translational Medicine 10(434), pii: eaaj1748.
32. Domenech de Celles M et al. (2014) Epidemiological evidence for herd immunity induced by acellular pertussis vaccines. Proceedings of the National Academy of Sciences of the United States of America 111, E16–E17.
33. Rohani P et al. (2010) Contact network structure explains the changing epidemiology of pertussis. Science 330, 982–985.
34. Domenech de Celles M et al. (2016) The pertussis enigma: reconciling epidemiology, immunology and evolution. Proceedings of the Royal Society B: Biological Sciences 283(1822), 20152309.
35. Edwards KM et al. (1995) Comparison of 13 acellular pertussis vaccines: overview and serologic response. Pediatrics 96, 548–557.
36. Commission CP (2015) Pharmacopoeia of People’s Republic of China 2015, Set 4, Chinese Edn. Beijing: China Medical Science and Technology Press.
37. Sun HY et al. (2001) Immunogenicity of two pertussis vaccine diphtheria and tetanus toxoid. Chinese Journal of Vaccines and Immunization 06, 37–40.
38. Gay NJ et al. (2000) Pertussis transmission in England and Wales. Lancet 355, 1553–1554.
39. Winter K et al. (2014) Pertussis epidemic – California, 2014. MMWR. Morbidity and Mortality Weekly Report 63, 1129–1132.
40. Sun M et al. (2010) Immunization status and risk factors of migrant children in densely populated areas of Beijing, China. Vaccine 28, 1264–1274.