Evaluation of the Brazilian surveillance system for adverse events following vaccination

Avaliação do sistema brasileiro de vigilância de eventos adversos pós-vacinação

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Objective: To describe and evaluate the Brazilian system of passive surveillance of adverse events following immunization (PSAEFI). Methods: The description and evaluation of PSAEFI were undertaken using the reported cases of adverse events following immunization with DTwP-Hib vaccine (AEFI-T), during the period from 2002 to 2005, using the Centers for Disease Control methodology. Results: The PSAEFI system, which provides national coverage, is designed to standardize practices in cases of adverse events following immunization (AEFI) and to identify highly reactogenic lots of vaccine. The PSAEFI system proved its usefulness, simplicity and flexibility; despite low sensitivity, overestimate the proportion of sever events, but it consistently described AEFI-T, identifying fever, convulsions and hypotonic-hyporesponsive episodes as the most common events. It showed that 49.7% of AEFI-T occur after the first dose, and that 72.8% occur within the first six hours after vaccination. It facilitates public health decisions and epidemiological investigations. It is timely, 46.1% of all AEFI-T being reported within 10 days after vaccination and its completeness ranges from 70 to 90%, depending on the item evaluated. Conclusions: The PSAEFI system proved useful for monitoring DTwP-Hib vaccine safety. We recommended the incorporation of new methodologies, such the use of sentinel cities/hospitals and computerized immunization registries in order to increase its sensitivity.

Keywords: immunization program; vaccination; product surveillance, postmarketing; epidemiologic surveillance; clinical trial, phase IV.
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Introduction

The Brazilian National Immunization Program (NIP), created in 1973, was a landmark in the history of public health in Brazil due to its high degree of organization and success in controlling diseases that, at that time, were responsible for high rates of infant morbidity and mortality.1,2

Like other countries, in view of concerns with vaccine safety and attempting to maintain high rates of adherence to the routine vaccination, Brazil implemented in 1998 a passive surveillance for adverse events following immunization. The Brazilian system of passive surveillance of adverse events following immunization (PSAEFI) is a nationwide surveillance system designed to monitoring vaccine safety and timely investigating suspected adverse events to support appropriate actions.5

The periodic evaluation of a surveillance system is a procedure that allows the degree of usefulness of the information provided by the surveillance to be determined, as well as allowing the event of interest to be estimated and the most vulnerable groups to be identified.7

The objective of this study was to describe the BSAEFI and evaluate its performance indicators during 2002 to 2005.

Resumo

Objetivos: Descrever e avaliar o Sistema brasileiro de vigilância passiva de eventos adversos pós-vacinação (SPVEAPV).

Métodos: A descrição e avaliação do SPVEAPV fundamentaram-se nas notificações de eventos adversos pós-vacina Tetravalente ou DTwP/Hib – vacina adsorvida difteria, tétano, pertussis e Haemophilus influenzae b (conjugada) - (EAPV-T), durante o período de 2002 a 2005. Empregou-se a metodologia proposta pelo Centers for Disease Control and Prevention. Resultados: O SPVEAPV apresenta abrangência nacional, tem por objetivos identificar e padronizar condutas frente a casos de eventos adversos pós-vacinação (EAPV) e identificar lotes reatogênicos. A vigilância é útil, simples e flexível, sua sensibilidade é baixa, superestima os eventos mais graves, mas descreve de forma consistente os EAPV-T, identificando a convulsão, a febre e o episódio hipotônico-hiporresponsivo como os mais frequentes, apontando a maior proporção de EAPV na primeira dose (49,7%) e nas primeiras seis horas após a vacinação (72,8%); é útil ao subsidiar decisões e investigações complementares; 46,1% das notificações são feitas até 10 dias após a vacinação; sua completude varia de 70% a 90%, conforme o item.

Conclusões: O SPVEAPV mostra-se útil no monitoramento da segurança da vacina DTwP/Hib, sendo, porém, recomendável a incorporação de novas metodologias como a de municípios e/ou hospitais sentinelas e a de sistemas informatizados de registros de imunização com a finalidade de elevar sua sensibilidade.

Palavras-chave: programas de imunização; vacinação; vigilância de produtos comercializados; vigilância epidemiológica; ensaio clínico fase IV.
Materials and Methods

This was a descriptive study in which the study area comprised the entire Brazilian territory including 26 states and one federal district, a total area of about 8.5 million km$^2$ and 190 million inhabitants$^{11}$. The study population included infants less than one year of age immunized with at least one dose of DTwP/Hib vaccine during 2002 to 2005. During the period of interest, the estimated population of infants less than one year of age in Brazil ranged from 3.3 million (in 2002) to 3.5 million (in 2005)$^{11}$.

The DTwP/Hib (or tetravalent) vaccine used in Brazil has been produced by Oswaldo Cruz Foundation (FIOCRUZ) Immunobiological Technology Institute (Bio-Manguinhos) in partnership with the Butantan Institute, which produces the DTwP fraction of the vaccine. The vaccine specifications are described elsewhere$^9$.

Information was obtained from the PSAEFI on reported cases of adverse events following DTwP/Hib vaccine (AEFI-T); the Brazilian Ministry of Health National Health System Database (DATASUS) on vaccine doses and coverage rate; and the Brazilian Institute of Geography and Statistics (IBGE) on population data. The A EFI-T case database is stored as part of the general PSAEFI database, denominated AEFI Information System (AEFI-IS).

The PSAEFI was evaluated according method proposed by the Centers for Disease Control and Prevention (CDC)$^7$. The magnitude and relevance of the event were described as well as the following surveillance components: objectives; case definition; sources of information; type of surveillance; analysis; and information dissemination. The same method was applied to assess qualitative and quantitative performance indicators.

The following concepts were used:
- Passive surveillance: information is obtained by spontaneous reporting;
- Active surveillance: information is obtained on a regular basis through direct contact between the surveillance team and information sources, or through electronic immunization records as part of electronic medical records$^8$.

Qualitative indicators:
- Usefulness: analyzes the degree to which the surveillance achieves its objectives, estimating the magnitude of AEFIs with the DTwP/Hib vaccine and whether it contributes to increasing the body of knowledge regarding the issue, identifying changing trends and stimulating research;
- Simplicity: evaluates the structure and its ease of operation;
- Flexibility: analyzes the ability of a surveillance to readily adapt to new information needs in response to changes in the nature or importance of the event of interest;
- Acceptability: evaluates compliance with the surveillance on the part of health professionals and health care facilities, one of the parameters being the completeness of the information included on the reporting form.

Quantitative indicators:
- Timeliness: evaluates the speed at which each step established by the surveillance system is achieved;
- Sensitivity: measures the proportion of true cases identified by the surveillance.

The consistency of the PSAEFI-T data was assessed based on case definition (presented and discussed in Results and Discussion). Only PSAEFI confirmed cases occurring among infants less than one year of age were included; cases associated with other vaccines, and those that were still under investigation were excluded. Since hypotonic-hypo-responsive episodes (HHEs) can be confused with convulsions, all AEFI-T involving HHEs accompanied by convulsion were registered as cases of convulsion alone$^{12}$.
A severe AEFI-T case was defined as any adverse event following DTwP/Hib vaccine that has resulted in death, hospitalization longer than 12 hours as well as cases of HHE, convulsions, encephalopathy, purpura, and hypersensitivity reaction within 2 hours after vaccination.\[13,14\]

For calculation of rates of reported cases the number of doses administered was used as the denominator. When there was more than one AEFI-T reported for the same child and dose, these events were considered as a single case with two or more events.

This study was approved by the Research Ethics Committee at Júlio Müller University Hospital at Universidade Federal de Mato Grosso (protocol no. 276/CEP-HUJM/06).

Results

Magnitude of the event and system description

Between 2002 and 2005, approximately 34 million doses of the DTwP/Hib vaccine were administered nationwide in Brazil. Administrative data show that, during the study period, the mean annual coverage among infants less than one year of age for the third dose of the DTwP/Hib vaccine was 85.0% nationwide (range, 66.0–100.0% among states), varying between 81.0% and 89.0% in the macroregions evaluated (north, south, northeast, central-west and southwest).\[15\]

There were 14,241 reported DTwP/Hib vaccine AEFI, corresponding to 11,558 infants, based on the case definition and inclusion/exclusion criteria adopted. The mean annual number of reported cases during the study period was 2,890, with a trend toward growth, increasing from 2,126 in 2002 to 3,383 in 2005.

The BSAEFI objectives included: i) to develop standards for identification of AEFI cases and management; ii) to analyze the national AEFI database; iii) to identify highly reactogenic batches and make decisions regarding management; iv) to ensure the population and health professionals that reliable immunobiologics are used in the NIP.\[5\]

Created by the NIP in 1998, the PSAEFI is a nationwide passive surveillance that receives reports from both public and private primary health care and hospital settings, especially from public ones. First restricted to the NIP, since 2008 the PSAEFI joint activities and promoting cooperation with the Brazilian Regulatory Health Agency (ANVISA) and the Oswaldo Cruz Foundation National Institute of Health Quality Control (INCQS).\[1\]

According to the PSAEFI a confirmed AEFI-T case is defined as any infant less than one year of age who following any dose of DTwP/Hib vaccine had one or more adverse events reported to the AEFI passive surveillance. Mild AEFI-T cases and those occurring more than 72 hours following immunization are excluded, with the exception of encephalopathy, which is considered to be an AEFI-T if identified within the first week after vaccination.\[5,16\]

The reports are made at the local level using a specific PSAEFI form. These forms are passed on to the state level on a monthly basis or immediately in the case of severe or uncommon events or outbreaks. Data are analyzed by trained technical staff at the state level. All confirmed cases are registered in the AEFI Information System (AEFI-IS) and passed on to the NIP on a monthly basis (Figure 1). The AEFI Information System was especially designed to store information on reported AEFI cases collected on standard reporting forms. It is used for recording and passing on reports from states to the NIP and for data analysis.

The NIP has, on a regular basis, updated the guidelines containing the technical norms employed for the training of personnel and the standardization of procedures.\[5,16\] One of those guidelines is designed exclusively for training local health care teams that work at vaccination sites.\[17\]
Evaluation of the system’s performance indicators

Data usefulness

The BSAEFI proved its usefulness. It was able to estimate the magnitude of the most common AEFI-T including seizure, fever and HHE (Table 1). It described their main features, showing a predominance of events following the first dose (49.7%) compared with the second (35.1%) and third doses (15.2%). It was found that 13.6% of these events occurred within the first hours following vaccination, 59.2% within one to six hours and only 2.3% occurred after 72 hours (Figure 2).

The weekly analysis of data showed peaks of reports, especially in macroregions of the country, where the surveillance proved to have higher sensitivity (Table 2). However, when vaccine batches were distributed, there were differences in the number of AEFI-T reported.

### Figure 1

![Surveillance system for adverse events following vaccination of the Brazilian National Immunization Program](source)

**Source:** Adapted from the Handbook of Adverse Events Following Immunization – Ministry of Health, Brazil, 2008.

**Fonte:** Adaptado de Manual de Eventos Adversos Pós-Vacinação – Ministério da Saúde, Brasil, 2008

*Routine flow: Adverse Events Following Immunization Reporting Form/AEFI-IS. Immediate flow: phone

*Fluxo de rotina: Ficha de Notificação e Investigação de Eventos Adversos Pós-Vacinação/FNIEAPV. Fluxo imediato: telefone

**Figure 1.** Surveillance system for adverse events following vaccination of the Brazilian National Immunization Program. Ministry of Health, Brazil

**Figura 1.** Sistema de vigilância de eventos adversos pós-vacinação do Programa Nacional de Imunizações. Ministério da Saúde, Brasil
Table 1. Number, percentage distribution and rate* of reported cases of major adverse events following diphtheria, tetanus, whole-cell pertussis-Haemophilus influenzae type b (DTwP-Hib) vaccination. Brazil, 2002–2005

<table>
<thead>
<tr>
<th>Adverse event following vaccination</th>
<th>Number of events (%)</th>
<th>Rate per 100,000 doses administered</th>
</tr>
</thead>
<tbody>
<tr>
<td>HHE</td>
<td>5208 (36.6)</td>
<td>15.3</td>
</tr>
<tr>
<td>Fever</td>
<td>4052 (28.4)</td>
<td>11.9</td>
</tr>
<tr>
<td>Fever&gt;39.5</td>
<td>2822 (19.8)</td>
<td>8.3</td>
</tr>
<tr>
<td>Fever&lt;39.5</td>
<td>1230 (8.6)</td>
<td>3.6</td>
</tr>
<tr>
<td>Convulsions</td>
<td>1992 (14.0)</td>
<td>5.9</td>
</tr>
<tr>
<td>Febrile convulsion</td>
<td>1578 (11.1)</td>
<td>4.6</td>
</tr>
<tr>
<td>Afebrile convulsion</td>
<td>414 (2.9)</td>
<td>1.2</td>
</tr>
<tr>
<td>Pain, redness and heat</td>
<td>462 (3.2)</td>
<td>1.4</td>
</tr>
<tr>
<td>Other severe and/or uncommon events</td>
<td>356 (2.5)</td>
<td>1.0</td>
</tr>
<tr>
<td>Abscess</td>
<td>333 (2.3)</td>
<td>1.0</td>
</tr>
<tr>
<td>Hypersensitivity&gt;2 hours</td>
<td>266 (1.9)</td>
<td>0.8</td>
</tr>
<tr>
<td>Other local reactions</td>
<td>248 (1.7)</td>
<td>0.7</td>
</tr>
<tr>
<td>Lump</td>
<td>246 (1.7)</td>
<td>0.7</td>
</tr>
<tr>
<td>Generalized rash</td>
<td>196 (1.3)</td>
<td>0.6</td>
</tr>
<tr>
<td>Induration</td>
<td>156 (1.1)</td>
<td>0.5</td>
</tr>
<tr>
<td>Hypersensitivity&lt;2 hours</td>
<td>133 (0.9)</td>
<td>0.4</td>
</tr>
<tr>
<td>Headache and vomiting</td>
<td>115 (0.8)</td>
<td>0.3</td>
</tr>
<tr>
<td>Anaphylaxis / anaphylactoid shock</td>
<td>26 (0.2)</td>
<td>0.1</td>
</tr>
<tr>
<td>Purpura</td>
<td>7 (0.05)</td>
<td>0.0</td>
</tr>
<tr>
<td>Acute encephalopathy</td>
<td>2 (0.01)</td>
<td>0.0</td>
</tr>
<tr>
<td>Total events</td>
<td>14,241 (100.0)</td>
<td>41.9</td>
</tr>
</tbody>
</table>

*per 100,000 doses administered; EHH: hypotonic-hyporesponsive episodes

with a higher number of reports associated were individually analyzed, no association was found with these peaks as reports were distributed over time. The individual analysis showed that batches with the highest number of reports associated had a proportion of serious adverse events similar to the average occurrence in the remaining batches (HHE: 25% to 55.9%; fever: 11.8 to 45.7%; and convulsions: 0% to 20.6%). Another study\(^{13}\) has pointed out the difficulty to identify highly reactogenic batches making it necessary to devise new monitoring strategies.

The PSAEFI databases provide information that can be used in research studies to further explore these adverse events\(^{18,19}\) and can provide input for the development and regular update of standards and regulations for safe vaccine use in Brazil.
**Table 2.** Proportion of municipalities with reported cases, number and proportion of severe cases of adverse events following diphtheria, tetanus, whole-cell pertussis-Haemophilus influenzae type b (DTwP-Hib) vaccination according to macroregion. Brazil, 2002–2005

<table>
<thead>
<tr>
<th>Macroregion</th>
<th>Infant Total</th>
<th>Municipalities Total</th>
<th>Number of events (%)</th>
<th>Reporting rate*</th>
<th>Severe cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>North</td>
<td>1,396,594</td>
<td>449</td>
<td>138 (30.7)</td>
<td>991 (7.0)</td>
<td>503 (61.9)</td>
</tr>
<tr>
<td>Northeast</td>
<td>4,158,867</td>
<td>1,792</td>
<td>462 (25.8)</td>
<td>1,957 (13.7)</td>
<td>1,057 (65.2)</td>
</tr>
<tr>
<td>South</td>
<td>1,832,141</td>
<td>1,189</td>
<td>538 (45.2)</td>
<td>4,127 (29.0)</td>
<td>1,631 (51.0)</td>
</tr>
<tr>
<td>Southeast</td>
<td>5,215,959</td>
<td>1,668</td>
<td>724 (43.4)</td>
<td>5,755 (40.4)</td>
<td>3,636 (74.5)</td>
</tr>
<tr>
<td>Midwest</td>
<td>960,640</td>
<td>466</td>
<td>175 (37.6)</td>
<td>1,411 (9.9)</td>
<td>715 (68.6)</td>
</tr>
<tr>
<td>Brazil</td>
<td>13,564,201</td>
<td>5,564</td>
<td>2,037 (36.6)</td>
<td>14,241 (100.0)</td>
<td>7,542 (65.3)</td>
</tr>
</tbody>
</table>

Source: Brazilian Ministry of Health, National Immunization Program, 2006

*Rate of adverse events following diphtheria, tetanus, whole-cell pertussis-Haemophilus influenzae type b (DTwP-Hib) immunization reported per 100,000 doses administered

**Simplicity**

The PSAEFI system is simple, the information flow being of low complexity, not involving laboratory tests or other specialized services (Figure 1\(^7\)). Another contributor to this simplicity is the fact that the public network of primary health care facilities not only functions as the principal source of reporting but is also responsible for the overwhelming majority of DTwP/Hib vaccinations carried out nationwide, as well
as for most of the medical appointments related to mild or medium-severity AEFIs with the DTwP-Hib vaccine. However, there are aspects that confer a certain degree of complexity upon the system. The data analysis and case confirmation require the involvement of well-trained teams. In addition, the perfect characterization of an AEFI is elusive, whether due to the absence of pathognomonic signs or to the unavailability of laboratory tests that would allow confirmation of the diagnosis. Nevertheless, the identification of a sign that could suggest an as-yet-unidentified adverse event requires the involvement of teams trained to carry out more complex investigations.

**Flexibility**

The BSAEFI proved to be flexible, since the introduction of the DTwP/Hib vaccine, as a replacement for the DTwP vaccine, did not alter the costs, routines, data sources or performance of the system.

**Acceptability**

Of a total of 5,564 municipalities, 1,999 (35.9%) reported an AEFI at least once during the study period (Table 2), and of them only 230 (4.1%) did it at least once a year. However, only 11 out of 5,564 municipalities (0.2%) have a population of more than 20,000 infants less than one year of age, which reduces the likelihood of identifying rare events.

Since it is mandatory the completion of the fields related to patient identification (name, age, gender), and date of vaccine administration and reporting in order to register the event in the AEFI database, this information was complete in 100% of reports.

Information related to the evolution of the AEFI-T, the strategy adopted in relation to maintaining or suspending the vaccine schedule, the type of hospitalization, the name of the laboratory that produced the vaccine and the vaccine lot number was included, respectively, in 79.5%, 75.4%, 69.6%, 92.3% and 84.7% of the reports.

**Timeliness**

Based on nationwide averages, in 25.3% of cases (range: 21.8% to 32.5%) the time between vaccine administration and reporting was one day and in 46.1% (range: 41.5% to 53.3%) was up to 10 days. Nationwide, an AEFI was registered in the AEFI database on the same date, 15 and 30 days after the reporting in 17%, 20% and 27% of cases, respectively, but in no less than 60 days in 57% of cases.

**Sensitivity**

The nationwide average AEFI-T reporting rate was 41.9 per 100,000 doses administered (ranging from 6 to 141 in the different geographical areas studied). The nationwide average rates for HHE and convulsions were 15 per 100,000 doses (ranging from 40 to 70), and 6 per 100,000 doses (ranging from 1 to 17). A comparison of these average rates found in the present study with other well-known studies showed that they are well below the expected, suggesting that the BSAEF has low sensitivity. This finding is corroborated by a recent study that estimated an average sensitivity of the BSAEFI-T for HHE and convulsions of 22.3% and 31.6% respectively; however, this same study reported widely varying rates, with sensitivity greater than 90% in some areas.

**Discussion**

The study showed that the PSAEFI has an adequate performance despite its low sensitivity. The surveillance proved useful in identifying and consistently describing AEFI-T most often associated with vaccines containing the pertussis component.

The surveillance’s good performance can be partly explained by the fact that Brazil has a broad network of primary health care facilities and a previously consolidated immunization program, with
trained, experienced teams and up-to-date guidelines for training courses that are also offered to health professionals working in local health care units\(^1,5,16\).

The study results showed that the PSAEFI has an appreciable degree of timeliness. Yet in approximately 50% of the AEFI-T, the time for register in the AEFI database was no less than 60 days. Despite this limitation the PSAEFI has successfully identified highly reactogenic vaccine batches\(^23,24\). In addition, it has good acceptability\(^6\), though it may be overestimated given the requirements for to register the event in the AEFI electronic database.

The average sensitivity of the BSAEFI was low. Sensitivity rates varied among states and were higher in those with better socioeconomic indicators and quality health services. Higher sensitivity rates were not associated with lower vaccination coverage\(^14\), i.e., the fact that health professionals are more aware of AEFI occurrence has not negatively affect adherence to vaccination.

As seen in other countries\(^22\), increasing AEFI reporting rates over the study period may be a result of increased sensitivity of the PSAEFI due to system improvements and due to the growing perception of the risk of AEFIs in certain social strata of the society. A similar trend was also reported by Freitas\(^13\) in the state of São Paulo, southeastern Brazil, over a longer study period while analyzing adverse events following DPT vaccination before the introduction of DTwP/Hib vaccine.

AEFI case definition may also affect sensitivity of the BSAEFI. The case definition used in Brazil focuses mainly on more serious events with public health relevance such as HHE and convulsions without regard for mild events such as fever. It has the advantage, especially in developing countries, of making surveillance less complex and costly as it examines smaller sets of data\(^13\). However, it may overestimate the relative importance of more serious events\(^3,4\) evidenced by fewer reports of fever than HHE while possibly underestimating late events, except for encephalopathy\(^5\).

The fact that the PSAEFI system is, in various aspects, a complex system of surveillance can produce favorable results, since it is a factor that induces the development of specialized human resources and the improvement of continuing education programs. In addition, this complexity can create conditions favorable to the strengthening of ties between the PSAEFI system and research groups, facilitating rapid responses in situations that elicit concern and anxiety on the part of the population in relation to vaccine safety\(^9\).

Many of the limitations identified in the Brazilian experience with the PSAEFI system are inherent to the use of passive surveillance, independent of the characteristics of the health care services in the country in which it is developed\(^3,4,7\). Perhaps the most important of such limitations is the low specificity, since the system frequently identified AEFIs temporarily associated with vaccination but without a causal relationship having been established\(^7,25\). Another limitation is its low sensitivity, as previously discussed\(^7,3\).

The surveillance’s limitations are also related to exposure to multiple vaccinations especially among infants less than one year of age; high number of potential AEFI\(^4\) as well as difficulty of identifying re-exposure to the same vaccine among individuals with previous reports of AEFI\(^1\). The choice for reporting more severe cases not only reduces the sensitivity of surveillance but also its representativeness\(^26\).

The favorable performance that the NIP has achieved in Brazil is the result of a set of public policies that strengthen the three pillars that on which the program rests: broad coverage, equity in access and vaccine safety\(^27\). These public policies have strengthened national vaccine manufacturers as well as quality control of vaccines, ensuring regular supply of biological products\(^27\). The NIP has promoted scientific and technological development, including training of multidisciplinary research
teams and continuing education\textsuperscript{9,28,29,30}. It has supported the conduction of periodic population-based surveys to monitor vaccine coverage and equitable access to vaccines\textsuperscript{31}.

The study showed the usefulness of passive surveillance systems for AEFI despite their limitations and challenges faced by developing countries like Brazil with large territorial areas and wide regional differences. These data support passive surveillance for AEFI as a strategic component for high levels of adherence to vaccination as it is a key tool for ensuring vaccine safety and reliability of immunization programs\textsuperscript{32}. They also indicate a need for new strategies to more effectively identify highly reactogenic vaccine batches as well as new methodologies such as sentinel hospitals/cities and electronic immunization records for increasing the system's sensitivity\textsuperscript{33}.

References


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