The Vaccine Adverse Event Reporting System (VAERS)

VAERS is a national post-licensure vaccine safety monitoring program co-managed by the Centers for Disease Control and Prevention (CDC) and the U.S. Food and Drug Administration (FDA). VAERS collects and analyzes information from reports of adverse events following receipt of U.S.-licensed vaccines. In recent years, VAERS has received approximately 40,000 U.S. reports annually, most of which describe mild adverse events like fever and injection site reactions. Very rarely, people experience serious adverse events following immunization. By monitoring such events, VAERS can help to identify important new safety concerns.

VAERS is a spontaneous reporting system, meaning that reports about adverse events can be submitted voluntarily by anyone. VAERS has limitations; data may, and often do, include incorrect and incomplete information. Underreporting and failure to report events occurs as well. Serious medical events are more likely to be reported than minor ones. Importantly, VAERS generally cannot determine cause and effect. A report to VAERS does not indicate that a vaccine caused an adverse event, only that the adverse event occurred sometime after vaccination. VAERS accepts all reports without judging the clinical seriousness of the adverse event or whether it was caused by the vaccine. More information on VAERS data can be found at: https://vaers.hhs.gov/data/dataguide.html

WHO CAN REPORT? Anyone can submit a VAERS report. Most reports are sent in by vaccine manufacturers and health care providers, but vaccine recipients, parents, and others may also submit reports.

WHAT SHOULD BE REPORTED? VAERS encourages reporting of any clinically significant adverse event that occurs after the administration of any vaccine licensed in the United States.

The National Childhood Vaccine Injury Act of 1986 requires health care providers to report:
- Any health event listed by the vaccine manufacturer as a contraindication to subsequent doses of the vaccine
- Any event listed in the Reportable Events Table that occurs within the specified time period after the vaccination.

A copy of the Reportable Events Table can be found on the following page, or at https://vaers.hhs.gov/docs/VAERS_Table_of_Reportable_Events_Following_Vaccination.pdf

HOW TO REPORT? There are two ways to report to VAERS:

- **Online.** Submit a VAERS report using the online reporting tool at https://vaers.hhs.gov/esub/index.jsp. Before you begin, review the Checklist for Completing the VAERS form at https://vaers.hhs.gov/reportevent.html. Information submitted using the online reporting tool is transmitted securely to VAERS.

- **Writable PDF Form.** Download the writable PDF form (located at https://vaers.hhs.gov/uploadFile/index.jsp) to your computer, complete it and then return to the VAERS website to upload the completed form. It is important that you use a desktop or laptop computer on which you can securely save a document that contains protected health information, personal identifiers or other sensitive personal or patient information. When you upload the form, the information is transmitted securely to VAERS.

  If you need further assistance with reporting to VAERS, please email info@vaers.org or call 1-800-822-7967. Operators are on duty from 9:00 a.m. to 5:00 p.m., Eastern Time, Monday through Friday.

For more information, visit the VAERS website at https://vaers.hhs.gov/
VAERS Table of Reportable Events Following Vaccination*

<table>
<thead>
<tr>
<th>Vaccine/Toxoid</th>
<th>Event and Interval from Vaccination</th>
</tr>
</thead>
</table>
| Tetanus in any combination: DTaP, DTP, DTP-Hib, DT, Td, TT, Tdap, DTaP-IPV, DTaP-IPV/Hib, DTaP-HepB-IPV | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Brachial neuritis (28 days)  
C. Shoulder Injury Related to Vaccine Administration (7 days)  
D. Any acute complications or sequelae (including death) of above events (interval - not applicable)  
E. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Pertussis in any combination: DTaP, DTP, DTP-Hib, Tdap, DTaP-IPV, DTaP-IPV/Hib, DTaP-HepB-IPV     | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Encephalopathy or encephalitis (7 days)  
C. Shoulder Injury Related to Vaccine Administration (7 days)  
D. Vasovagal syncope (7 days)  
E. Any acute complications or sequelae (including death) of above events (interval - not applicable)  
F. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Measles, mumps and rubella in any combination: MMR, MMRV, MM                                      | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Encephalopathy or encephalitis (15 days)  
C. Shoulder Injury Related to Vaccine Administration (7 days)  
D. Vasovagal syncope (7 days)  
E. Any acute complications or sequelae (including death) of above events (interval - not applicable)  
F. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Rubella in any combination: MMR, MMRV                                                           | A. Chronic arthritis (42 days)  
B. Any acute complications or sequelae (including death) of above event (interval - not applicable)  
C. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Measles in any combination: MMR, MMRV, MM                                                         | A. Thrombocytopenic purpura (7-30 days)  
B. Vaccine-strain measles viral infection in an immunodeficient recipient  
  • Vaccine-strain virus identified (interval – not applicable)  
  • If strain determination is not done or if laboratory testing is inconclusive (12 months)  
C. Any acute complications or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Oral Polio (OPV)                                                                                   | A. Paralytic polio  
  • in a non-immunodeficient recipient (30 days)  
  • in an immunodeficient recipient (6 months)  
  • in a vaccine-associated community case (interval - not applicable)  
B. Vaccine-strain polio viral infection  
  • in a non-immunodeficient recipient (30 days)  
  • in an immunodeficient recipient (6 months)  
  • in a vaccine-associated community case (interval - not applicable)  
C. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Inactivated Polio in any combination: IPV, DTaP-IPV, DTaP-IPV/Hib, DTaP-HepB-IPV                  | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Shoulder Injury Related to Vaccine Administration (7 days)  
C. Vasovagal syncope (7 days)  
D. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
E. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| Hepatitis B in any combination: HepB, HepA-HepB, DTaP-HepB-IPV, Hib-HepB                            | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Shoulder Injury Related to Vaccine Administration (7 days)  
C. Vasovagal syncope (7 days)  
D. Any acute complications or sequelae (including death) of above events (interval - not applicable)  
E. Events described in manufacturer’s package insert as contraindications to additional doses of vaccine (interval - see package insert) |

*Updated July 12, 2017
<table>
<thead>
<tr>
<th>Vaccine/Toxoid</th>
<th>Event and Interval from Vaccination</th>
</tr>
</thead>
</table>
| **Haemophilus influenzae type b in any combination (conjugate): Hib, Hib-HepB, DTaP-IPV/Hib, Hib-MenCY** | A. Shoulder Injury Related to Vaccine Administration (7 days)  
B. Vasovagal syncope (7 days)  
C. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Varicella in any combination: VAR, MMRV** | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Disseminated varicella vaccine-strain viral disease  
- Vaccine-strain virus identified (time interval unlimited)  
- If strain determination is not done or if laboratory testing is inconclusive (42 days)  
C. Varicella vaccine-strain viral reactivation (time interval unlimited)  
D. Shoulder Injury Related to Vaccine Administration (7 days)  
E. Vasovagal syncope (7 days)  
F. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
G. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Rotavirus (monovalent or pentavalent) RV1, RV5** | A. Intussusception (21 days)  
B. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
C. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Pneumococcal conjugate (7-valent or 13-valent) PCV7, PCV13** | A. Shoulder Injury Related to Vaccine Administration (7 days)  
B. Vasovagal syncope (7 days)  
C. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Hepatitis A in any combination: HepA, HepA-HepB** | A. Shoulder Injury Related to Vaccine Administration (7 days)  
B. Vasovagal syncope (7 days)  
C. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Seasonal influenza (trivalent inactivated influenza, quadrivalent inactivated influenza, live attenuated influenza): IIIV3, IIV4, RIV3, cIIIV3, LAIV4** | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Shoulder Injury Related to Vaccine Administration (7 days)  
C. Vasovagal syncope (7 days)  
D. Guillain-Barré Syndrome (42 days)  
E. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
F. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Meningococcal: MCV4, MPSV4, Hib-MenCY, MenACWY, MenB** | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Shoulder Injury Related to Vaccine Administration (7 days)  
C. Vasovagal syncope (7 days)  
D. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
E. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Human Papillomavirus (quadrivalent, bivalent, or 9 valent): 9vHPV, 4vHPV, 2vHPV** | A. Anaphylaxis or anaphylactic shock (7 days)  
B. Shoulder Injury Related to Vaccine Administration (7 days)  
C. Vasovagal syncope (7 days)  
D. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
E. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
| **Any new vaccine recommended by the Centers for Disease Control and Prevention for routine administration to children.** | A. Shoulder Injury Related to Vaccine Administration (7 days)  
B. Vasovagal syncope (7 days)  
C. Any acute complication or sequelae (including death) of above events (interval - not applicable)  
D. Events described in manufacturer's package insert as contraindications to additional doses of vaccine (interval - see package insert) |
The Reportable Events Table (RET) reflects what is reportable by law (42 USC 300aa-25) to the Vaccine Adverse Event Reporting System (VAERS) including conditions found in the manufacturer package insert. In addition, healthcare professionals are encouraged to report any clinically significant or unexpected events (even if not certain the vaccine caused the event) for any vaccine, whether or not it is listed on the RET. Manufacturers are also required by regulation (21 CFR 600.80) to report to the VAERS program all adverse events made known to them for any vaccine.

Note that the RET differs from the Vaccine Injury Table (VIT) regarding timeframes of adverse events. Timeframes listed on the RET reflect what is required for reporting, but not what is required for compensation. To view timeframes for compensation, please see the VIT at https://www.hrsa.gov/vaccinecompensation/vaccineinjurytable.pdf

**Represents the onset interval between vaccination and the adverse event. For a detailed explanation of terms, see the Vaccine Injury Table at https://www.hrsa.gov/vaccinecompensation/vaccineinjurytable.pdf
The Vaccine Injury Compensation Program (VICP)

The VICP is a no-fault alternative to the traditional tort system for resolving vaccine injury claims. It was established as part of the National Childhood Vaccine Injury Act of 1986, after a rash of lawsuits against vaccine manufacturers and healthcare providers threatened to cause vaccine shortages and reduce vaccination rates.

The VICP is administered jointly by the U.S. Department of Health and Human Services (HHS), the U.S. Court of Federal Claims (the Court), and the U.S. Department of Justice (DOJ). The VICP is located in the HHS, Health Resources and Services Administration (HRSA), Healthcare Systems Bureau, Division of Vaccine Injury Compensation.

Briefly, an individual claiming a vaccine-related injury or death files a petition for compensation with the Court, and may be represented by an attorney. A HHS physician reviews the petition to determine whether it meets the medical criteria for compensation. A recommendation is provided to the Court. The HHS position is presented before a “special master,” who makes the decision for compensation under the VICP. A decision may be appealed to a judge of the Court, then to the Federal Circuit Court of Appeals, and eventually to the U.S. Supreme Court.

A petitioner may file a claim in civil court against the vaccine company and/or the vaccine administrator only after first filing a claim under the VICP and then rejecting the decision of the Court.

Who Can File a Claim?

- You may file a claim if you received a vaccine covered by the VICP and believe that you have been injured by this vaccine.
- You may also file a claim if you are a parent or legal guardian of a child or disabled adult who received a vaccine covered by the VICP and believe that the person was injured by this vaccine.
- You may file a claim if you are the legal representative of the estate of a deceased person who received a vaccine covered by the VICP and believe that the person’s death resulted from the vaccine injury.
- You may file a claim if you are not a United States citizen.
- Some people who receive vaccines outside of the U.S. may be eligible for compensation. See the VICP website for more details.
- In addition, to be eligible to file a claim, the effects of the person’s injury must have:
  1. lasted for more than 6 months after the vaccine was given; or
  2. resulted in a hospital stay and surgery; or
  3. resulted in death.

There is no age restriction on who may file a claim. Anyone receiving a vaccine covered by the VICP, no matter their age, can file a claim or have one filed on their behalf.

To learn how to file a claim, see the VICP website at http://www.hrsa.gov/vaccinecompensation/fileclaim.html.

Vaccines covered by VICP are diphtheria, tetanus, pertussis, Hib, hepatitis A, hepatitis B, human papillomavirus, seasonal influenza, measles, mumps, rubella, meningococcal, polio, pneumococcal conjugate, rotavirus, and varicella, in any combination. (Additional vaccines may be added in the future.)
The **Vaccine Injury Table** makes it easier for some people to get compensation. The Table lists and explains injuries and conditions that are presumed to be caused by vaccines. It also lists time periods in which the first symptom of these injuries and conditions must occur after receiving the vaccine. If the first symptom of these injuries/conditions occurs within the listed time periods, it is presumed that the vaccine was the cause of the injury or condition unless another cause is found. For example, if a patient received the tetanus vaccine and had a severe allergic reaction (anaphylaxis) within 4 hours after receiving the vaccine, then it is presumed that the tetanus vaccine caused the injury, if no other cause is found.

If an injury or condition is not on the Table or if it did not occur within the time period on the Table, the petitioner must prove that the vaccine caused the injury or condition. Such proof must be based on medical records or opinion, which may include expert witness testimony.

A copy of the Vaccine Injury Table is on the following page, or can be found online at [http://www.hrsa.gov/vaccinecompensation/vaccinetable.html](http://www.hrsa.gov/vaccinecompensation/vaccinetable.html). A comprehensive explanation of terms used in the table accompanies the online version.

March 2013 (Revised March 2017)

For more information, visit the VICP website at [http://www.hrsa.gov/vaccinecompensation](http://www.hrsa.gov/vaccinecompensation).
# National Childhood Vaccine Injury Act: Vaccine Injury Table

This table, supplemented with definitions and other explanatory material, can be found on the National Vaccine Injury Compensation Program’s website at [www.hrsa.gov/vaccinecompensation/vaccineinjurytable.pdf](http://www.hrsa.gov/vaccinecompensation/vaccineinjurytable.pdf).

<table>
<thead>
<tr>
<th>Vaccine</th>
<th>Illness, disability, injury or condition covered</th>
<th>Time period for first symptom or manifestation of onset or of significant aggravation after vaccine administration</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Vaccines containing tetanus toxoid (e.g., DTaP, DTP, DT, Td, or TT)</td>
<td>A. Anaphylaxis</td>
<td>≤4 hours</td>
</tr>
<tr>
<td></td>
<td>B. Brachial Neuritis</td>
<td>2-28 days (not less than 2 days and not more than 28 days)</td>
</tr>
<tr>
<td></td>
<td>C. Shoulder Injury Related to Vaccine Administration</td>
<td>≤48 hours</td>
</tr>
<tr>
<td></td>
<td>D. Vasovagal syncope</td>
<td>≤1 hour</td>
</tr>
<tr>
<td>II. Vaccines containing whole cell pertussis bacteria, extracted or partial cell pertussis bacteria, or specific pertussis antigen(s) (e.g., DTP, DTaP, P, DTP-Hib)</td>
<td>A. Anaphylaxis</td>
<td>≤4 hours</td>
</tr>
<tr>
<td></td>
<td>B. Encephalopathy or encephalitis</td>
<td>≤72 hours</td>
</tr>
<tr>
<td></td>
<td>C. Shoulder Injury Related to Vaccine Administration</td>
<td>≤48 hours</td>
</tr>
<tr>
<td></td>
<td>D. Vasovagal syncope</td>
<td>≤1 hour</td>
</tr>
<tr>
<td>III. Vaccines containing measles, mumps, and rubella virus or any of its components (e.g., MMR, MM, MMRV)</td>
<td>A. Anaphylaxis</td>
<td>≤4 hours</td>
</tr>
<tr>
<td></td>
<td>B. Encephalopathy or encephalitis</td>
<td>5-15 days (not less than 5 days and not more than 15 days)</td>
</tr>
<tr>
<td></td>
<td>C. Shoulder Injury Related to Vaccine Administration</td>
<td>≤48 hours</td>
</tr>
<tr>
<td></td>
<td>D. Vasovagal syncope</td>
<td>≤1 hour</td>
</tr>
<tr>
<td>IV. Vaccines containing rubella virus (e.g., MMR, MMRV)</td>
<td>A. Chronic arthritis</td>
<td>7-42 days (not less than 7 days and not more than 42 days)</td>
</tr>
<tr>
<td>V. Vaccines containing measles virus (e.g., MMR, MM, MMRV)</td>
<td>A. Thrombocytopenic purpura</td>
<td>7-30 days (not less than 7 days and not more than 30 days)</td>
</tr>
<tr>
<td></td>
<td>B. Vaccine-Strain Measles Viral Infection in an immunodeficient recipient:</td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td>- Vaccine-strain virus identified</td>
<td>≤12 months</td>
</tr>
<tr>
<td></td>
<td>- If strain determination is not done or if laboratory testing is inconclusive</td>
<td>≤30 days</td>
</tr>
<tr>
<td></td>
<td>- in a non-immunodeficient recipient</td>
<td>≤6 months</td>
</tr>
<tr>
<td></td>
<td>- in an immunodeficient recipient</td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td>- in a vaccine associated community case</td>
<td>≤30 days</td>
</tr>
<tr>
<td>VI. Vaccines containing polio live virus (OPV)</td>
<td>A. Paralytic Polio</td>
<td>≤6 months</td>
</tr>
<tr>
<td></td>
<td>- in a non-immunodeficient recipient</td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td>- in an immunodeficient recipient</td>
<td>≤30 days</td>
</tr>
<tr>
<td></td>
<td>- in a vaccine associated community case</td>
<td>≤6 months</td>
</tr>
<tr>
<td></td>
<td>B. Vaccine-Strain Polio Viral Infection</td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td>- in a non-immunodeficient recipient</td>
<td>≤30 days</td>
</tr>
<tr>
<td></td>
<td>- in an immunodeficient recipient</td>
<td>≤6 months</td>
</tr>
<tr>
<td></td>
<td>- in a vaccine associated community case</td>
<td>Not applicable</td>
</tr>
<tr>
<td>VII. Vaccines containing polio inactivated virus (e.g., IPV)</td>
<td>A. Anaphylaxis</td>
<td>≤4 hours</td>
</tr>
<tr>
<td></td>
<td>B. Shoulder Injury Related to Vaccine Administration</td>
<td>≤48 hours</td>
</tr>
<tr>
<td></td>
<td>C. Vasovagal syncope</td>
<td>≤1 hour</td>
</tr>
<tr>
<td>VIII. Hepatitis B vaccines</td>
<td>A. Anaphylaxis</td>
<td>≤4 hours</td>
</tr>
<tr>
<td></td>
<td>B. Shoulder Injury Related to Vaccine Administration</td>
<td>≤48 hours</td>
</tr>
<tr>
<td></td>
<td>C. Vasovagal syncope</td>
<td>≤1 hour</td>
</tr>
<tr>
<td>Vaccine</td>
<td>Illness, disability, injury or condition covered</td>
<td>Time period for first symptom or manifestation of onset or of significant aggravation after vaccine administration</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| IX. *Haemophilus influenzae* type b (Hib) vaccines | A. Shoulder Injury Related to Vaccine Administration  
B. Vasovagal syncope                                                                                           | ≤48 hours  
≤1 hour                                                                                                            |
| X. Varicella vaccines                       | A. Anaphylaxis  
B. Disseminated varicella vaccine-strain viral disease:  
- Vaccine-strain virus identified  
- If strain determination is not done or if laboratory testing is inconclusive  
C. Varicella vaccine-strain viral reactivation  
D. Shoulder Injury Related to Vaccine Administration  
E. Vasovagal syncope                                                                                           | <4 hours  
Not applicable  
7-42 days (not less than 7 days and not more than 42 days)  
Not applicable  
≤48 hours  
≤1 hour                                                                                                            |
| XI. Rotavirus vaccine                       | A. Intussusception                                                                                                  | 1-21 days (not less than 1 day and not more than 21 days)                                                   |
| XII. Pneumococcal conjugate vaccines        | A. Shoulder Injury Related to Vaccine Administration  
B. Vasovagal syncope                                                                                           | ≤48 hours  
≤1 hour                                                                                                            |
| XIII. Hepatitis A vaccines                 | A. Shoulder Injury Related to Vaccine Administration  
B. Vasovagal syncope                                                                                           | ≤48 hours  
≤1 hour                                                                                                            |
| XIV. Seasonal influenza vaccines            | A. Anaphylaxis  
B. Shoulder Injury Related to Vaccine Administration  
C. Vasovagal syncope  
D. Guillain-Barré Syndrome                                                                           | <4 hours  
≤48 hours  
≤1 hour  
3-42 days (not less than 3 days and not more than 42 days)                                                  |
| XV. Meningococcal vaccines                 | A. Anaphylaxis  
B. Shoulder Injury Related to Vaccine Administration  
C. Vasovagal syncope                                                                                           | ≤4 hours  
≤48 hours  
≤1 hour                                                                                                            |
| XVI. Human papillomavirus (HPV) vaccines   | A. Anaphylaxis  
B. Shoulder Injury Related to Vaccine Administration  
C. Vasovagal syncope                                                                                           | <4 hours  
≤48 hours  
≤1 hour                                                                                                            |
| XVII. Any new vaccine recommended by the Centers for Disease Control and Prevention for routine administration to children, after publication by the Secretary of a notice of coverage | A. Shoulder Injury Related to Vaccine Administration  
B. Vasovagal syncope                                                                                           | ≤48 hours  
≤1 hour                                                                                                            |

*(Applies Only to Petitions for Compensation Filed under the National Vaccine Injury Compensation Program on or after March 21, 2017)*
Countermeasures Injury Compensation Program (CICP)

Overview

The Countermeasures Injury Compensation Program (CICP) is a Federal program that provides benefits to individuals who are seriously injured as a result of the administration or use of a covered countermeasure. CICP also provides death benefits to certain survivors if death directly resulted from receipt of a covered countermeasure. Covered countermeasures may include vaccines, antivirals, drugs, biologics, or medical devices used to prevent, treat, or diagnose an illness that the Secretary of the United States Department of Health and Human Services (the Secretary) declares to be an actual or potential public health emergency. Examples of currently covered countermeasures are pandemic influenza vaccines including the 2009 pandemic H1N1 influenza vaccine, antivirals (e.g., Tamiflu®, Relenza®, peramivir), ventilation assistance devices (e.g., mechanical ventilators), and respiratory protection devices (e.g., N-95 masks) used to treat, diagnose or prevent pandemic influenza. In addition, countermeasures, including vaccines, used to diagnose, treat or prevent smallpox, anthrax, botulism, and acute radiation syndrome are currently covered. Adverse events during pre-licensure testing may be covered as well.

This Program was established by the Public Readiness and Emergency Preparedness Act of 2005 (PREP Act), 42 U.S.C. § 247d-6e. The PREP Act also confers broad liability protections covering the manufacture, testing, development, distribution, or use of the designated covered countermeasure.

Filing Deadline and Application and Review Process

Individuals have one (1) year from the date the vaccine or other covered countermeasure was administered or used to request compensation benefits. If their injury is added to a Countermeasures Injury Table, then they may also have one year from the effective date of the Table addition to file. To file a claim, individuals must submit a Request for Benefits Form and the Authorization for Use or Disclosure of Health Information Form to request medical records from each health care provider who treated the injured person. In addition, medical records from one year before the injury to the present time must be submitted. Health care providers should send medical records directly to the Program. All documents should be sent to:
Health Resources and Services Administration
Countermeasures Injury Compensation Program
5600 Fishers Lane, Room 11C-26
Rockville, MD 20857

After an individual has submitted a complete Request for Benefits package, CICP medical staff reviews it to determine if the individual is eligible for compensation. An individual may be eligible for compensation if compelling, reliable, valid, medical and scientific evidence exists demonstrating that the injury for which compensation is sought was caused by the administration or use of a covered countermeasure and no other more likely cause of the injury is found. If an individual is found eligible for compensation, the type and amount of benefits are determined by the Program. If an individual is not eligible for compensation, he/she may request the Associate Administrator of the Healthcare Systems Bureau, HRSA, to reconsider the Program’s decision. The Associate Administrator will convene an independent panel to review the Program’s decision, make its own findings, and make a recommendation. The Associate Administrator will review this recommendation and make a final decision.
Benefits Available

Eligible individuals may be compensated for reasonable and necessary unreimbursable medical expenses and for lost employment income at the time of the injury. Death benefits may be paid to certain survivors of covered countermeasures recipients who have died as a direct result of the covered countermeasure injury. The U.S. Department of Health and Human Services is the payer of last resort. Therefore, payments are reduced by those of other third party payers.

Contact Information
Website: http://www.hrsa.gov/cicp/
E-mail: CICP@hrsa.gov
Phone: 1-855-266-CICP (2427)

Updated March 18, 2013