

ma; Gammagard; Globuman†; Intraglobin F; Octagam; Pentaglobin; Venoglobulin-S†; **Hung.**: Gammagard; Gammanorm; Humaglobin; Intratect; Octagam; Pentaglobin; Vivaglobin; **India.**: Gamafine; **Indon.**: Gammune N; Gammaraas; **Irl.**: Intraglobin; **Israel.**: Beriglobin P; Endobulin†; Flebogamma; Gammagard†; Intraglobin F; Omri-IgG; Sandoglobulin; Venoglobulin; Vigam; **Ital.**: Biaven; Endobulin; Flebogamma; Gamma-Venin P†; Gammagard; Globuman†; Haimaven†; Ig Gamma†; Ig Vena; Intraglobin; Isiven; Pentaglobulin; **Malaysia.**: Flebogamma; Gammagard†; Globuman†; Intraglobin F†; IV-Globulin; Pentaglobin†; Venoglobulin-S†; Vigam; **Mex.**: Beriglobina P; Gamma-Venin P; Gammagard†; Isiven†; Octagam; Pentaglobin; Sandoglobulina; Seroglobulin†; Vigam; **Neth.**: Endobulin†; Flebogamma; Gammagard; Gammanorm; GammaQuin; Ivegam; Octagam; Subcuvia; Vivaglobin; **Norw.**: Gammaglobulin†; Gammanorm; Octagam; **NZ.**: Intragam; Octagam; Sandoglobulin; **Philipp.**: Gammune N; Gammagard; IV-Globulin S; **Pol.**: Endobulin; Gamma-Globulina; Gammagard; Intraglobin F; Intratect; Kioiv; Pentaglobin; Sandoglobulin; Subcuvia; Venimmun; **Port.**: Flebogamma; Gammagard; Gammanorm; Gamunex; Globuman†; Ig Vena; Octagam; Sandoglobulina; Subcuvia; Vivaglobin; **Rus.**: Gammune N (Гаммун Н); Humaglobin (Хумаглобин); Immuovenin (Иммуовенин); Octagam (Октагам); **S.Afr.**: Beriglobin; Endobulin†; Intragam; Intraglobin F; Pentaglobin; Polygam; **Singapore.**: Flebogamma; Gammagard†; Intraglobin F; Pentaglobin; Venoglobulin†; Vigam†; **Spain.**: Beriglobina P; Endobulin; Flebogamma; Gammagard; Gammaglobulina; Globuman†; Octagamocta; **Swed.**: Beriglobin; Endobulin; Gammagard; Gammanorm; Gammonativ; Octagam; Polyglobin†; Subcuvia; Xepol; **Switz.**: Beriglobin; Endobulin; Gammagard; Globuman; Intraglobin F; Octagam; Pentaglobin; Redimune; **Thai.**: Flebogamma; Gammaraas; Globuman†; Ig Vena; Intraglobin; Octagam; Pentaglobin; Venoglobulin-S†; Vigam; **Turk.**: Bisek; Endobulin; Flebogamma; Gammune N; Gammara; Globuman†; IG Vena NIV; Intraglobin; Isiven; Octagam; Pentaglobin; Subcuvia; Tegeline; Vigam; **UK.**: Flebogamma; Gammabulin†; Gammagard; Kioiv; Octagam; Sandoglobulin; Subcuvia; Subgam; Vigam; Vivaglobin; **USA.**: Carimune; Flebogamma; Gamastan; Gammune N†; Gammagard; Gammara-P†; Gamunex; Ivegam†; Octagam; Panglobulin†; Polygam; Privigen; Venoglobulin†; Vivaglobin; **Venez.**: Flebogamma; Sandoglobulin†; Venoglobulina†.

Multi-ingredient: **Arg.**: Biotaer Gamma†; Histaglobin; **Austria:** Histaglobin; **Chile:** Pentaglobin; **Cz.**: Histaglobin†; **Ger.**: Histadestal; **India:** Histaglobulin; **Pol.**: Histaglobulina; **S.Afr.**: Histaglobin.

Pertussis Immunoglobulins

Immunoglobulinas contra la tos ferina.

ATC — J06BB13.

Pharmacopoeias. Many pharmacopoeias, including *US*, have monographs.

USP 31 (Pertussis Immune Globulin). A sterile solution of globulins derived from the plasma of adult human donors who have been immunised with pertussis vaccine. It may contain glycine as a stabilising agent, and a suitable preservative. It should be stored at 2° to 8°.

Adverse Effects and Precautions

As for immunoglobulins in general, p.2201.

Interactions

As for immunoglobulins in general, p.2201.

Uses and Administration

Pertussis immunoglobulins have been used for passive immunisation against pertussis (whooping cough) and to prevent or modify pertussis in susceptible persons who have been exposed to infection.

Preparations

USP 31: Pertussis Immune Globulin.

Pertussis Vaccines

Vacunas de la tos ferina.

ATC — J07AJ01; J07AJ02.

Pharmacopoeias. Many pharmacopoeias, including *Eur.* (see p.vii), have monographs.

Ph. Eur. 6.2 (Pertussis Vaccine (Adsorbed); Vaccinum Pertussis Adsorbatum). A sterile suspension of inactivated whole cells of one or more strains of *Bordetella pertussis* in saline to which hydrated aluminium phosphate or aluminium hydroxide has been added. It may contain a suitable antimicrobial preservative. The estimated potency is not less than 4 units per dose. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light.

The BP 2008 states that wP may be used on the label.

Ph. Eur. 6.2 (Pertussis Vaccine (Acellular; Component, Adsorbed); Vaccinum Pertussis Sine Cellulis ex Elementis Praeparatum Adsorbatum). A preparation of individually prepared and purified antigenic components of *Bordetella pertussis* adsorbed on a mineral carrier such as aluminium hydroxide or hydrated aluminium phosphate. It contains either a suitably prepared pertussis toxinoid or a pertussis toxin-like protein free from toxic properties produced by expression of a genetically modified form of the corresponding gene. It may also contain filamentous haemagglutinin, pertactin, and other defined antigens such as fimbrial-2 and fimbrial-3 antigens. The final vaccine contains not more than 100 units of bacterial endotoxin per dose. It may contain a suitable antimicrobial preservative. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light. The BP 2008 states that aP may be used on the label.

Ph. Eur. 6.2 (Pertussis Vaccine (Acellular; Co-purified, Adsorbed); Vaccinum Pertussis Sine Cellulis Copurificatum Adsorbatum). A preparation of antigenic components of *Bordetella pertussis* adsorbed on a mineral carrier such as aluminium hydroxide or hydrated aluminium phosphate. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light. The BP 2008 states that aP may be used on the label.

Adverse Effects

As for vaccines in general, p.2201.

Local reactions may occur at the site of injection of pertussis vaccines or pertussis-containing vaccines and use may be followed by fever and irritability. Local reactions and fever occur less frequently with the acellular vaccine than with whole-cell vaccine, especially in children over 6 months of age and adults. However, booster doses of acellular pertussis-containing vaccines are associated with an increased risk of injection site reactions.

Severe reactions which have been reported include persistent screaming and generalised collapse but these effects were generally associated with an earlier type of vaccine and the reactions are stated to be rarely observed with the currently available vaccines.

Rare neurological adverse reactions have included convulsions and encephalopathy. There has been much debate, however, on the causal role of pertussis vaccine in such reactions (see below for detailed discussion). It should be remembered that neurological complications occur more frequently as a consequence of pertussis infection than from vaccination.

Asthma. A higher incidence of asthma was reported in 243 children who had received whole-cell pertussis vaccine than in 203 children who had not.¹ However, follow-up of a large Swedish study² showed no difference in the incidence of wheezing or allergic reactions between children who had received diphtheria, tetanus, and whole-cell pertussis vaccines and those who had not. A later prospective study³ also found no evidence that whole-cell pertussis vaccination increased the risk of wheezing illness in young children. Furthermore, no association was found between pertussis vaccination in infancy and development of asthma in children aged up to 7 years in a later study of the same group of children.⁴

1. Odent MR, *et al.* Pertussis vaccination and asthma: is there a link? *JAMA* 1994; **272**: 592–3.
2. Nilsson L, *et al.* Lack of association between pertussis vaccination and symptoms of asthma and allergy. *JAMA* 1996; **275**: 760.
3. Henderson J, *et al.* Pertussis vaccination and wheezing illnesses in young children: prospective cohort study. *BMJ* 1999; **318**: 1173–6.
4. Maitra A, *et al.* Pertussis vaccination in infancy and asthma or allergy in later childhood: birth cohort study. *BMJ* 2004; **328**: 925–6.

Effects on the nervous system. There has been continuing debate over several decades concerning the perceived link between pertussis vaccination and brain damage. Anxiety among both the public and health care professionals in the UK in the mid-1970s over the safety of whole-cell pertussis vaccines led to a fall in the acceptance rates for infant vaccination and major epidemics of pertussis in 1977/79 and 1981/83. Since that time confidence has been restored and, with the introduction of acellular vaccines, the vast majority of infants now receive the vaccine before their second birthday. Comparison of whole-cell vaccines with acellular vaccines has since confirmed that the latter are associated with a greatly reduced incidence of serious neurological disorders.¹

The consensus of opinion now seems to be that there is a temporal, but not necessarily causal, relationship between whole-cell pertussis vaccine and acute neurological illness that may occasionally lead to long-term dysfunction, and that risks of not immunising are greater than the potential risks associated with the vaccine.

The difficulty in ascertaining whether a causal relationship exists between whole-cell pertussis vaccine (usually given as diphtheria, tetanus, and pertussis (DTP) vaccine) and acute neurological reactions arises partly because primary vaccination is given at an age when neurological dysfunction with other causes is often first manifested. The observed temporal relationship may be entirely coincidental, may result from indirect factors such as pyrexia after vaccination, or may represent a direct effect of DTP vaccine. Much of the evidence is based on large epidemiological studies,^{2–7} in particular the National Childhood Encephalopathy Study (NCES)⁸ from the UK and its 10-year follow-up.⁹ Serious acute neurological illnesses reported to the NCES⁸ were found to be more common in infants immunised within 7 days (relative risk 2.4), and especially within 72 hours of onset, than in unimmunised children. For previously normal children, irrespective of outcome, the risk was estimated as 1 in 110 000 injections. In a subset of cases diagnosed as infantile spasms,¹⁰ no link with vaccination was found overall, but there was a small excess of cases of infantile spasm in previously normal children who had received either DTP or diphtheria and tetanus vaccines during the previous 7 days (relative risk 2.0–2.5) followed by a corresponding deficit during the next 3 weeks. This suggested that vaccination may trigger the onset of spasms in a child with an underlying predisposition.

In 1991, the USA Institute of Medicine reviewed the available data, including the NCES results, and concluded that a causal relationship between the whole-cell pertussis component of DTP vaccine and acute encephalopathy probably existed, with an estimated excess risk of 0 to 10.5 per million vaccinations.¹¹ They concurred with the conclusion that a causal relationship between vaccination and infantile spasm was unlikely.

The NCES 10-year follow-up found that children who had had a serious acute neurological illness (excluding infantile spasms) had an increased risk of death or long-term dysfunction, but the risk was no greater in children given DTP vaccine in the 7 days before the original acute illness.⁹ The National Vaccines Advisory Committee concluded that the results were insufficient to determine whether DTP vaccine influenced the development of chronic neurological dysfunction, and this conclusion has been accepted by both the Advisory Committee on Immunization Practices¹² and the American Academy of Pediatrics.¹³

1. Geier DA, Geier MR. An evaluation of serious neurological disorders following immunization: a comparison of whole-cell pertussis and acellular pertussis vaccines. *Brain Dev* 2004; **26**: 296–300.
2. Pollock TM, Morris J. A 7-year survey of disorders attributed to vaccination in North West Thames Region. *Lancet* 1983; **i**: 753–7.
3. Pollock TM, *et al.* Symptoms after primary immunisation with DTP and with DT vaccine. *Lancet* 1984; **ii**: 146–9.
4. Walker AM, *et al.* Neurologic events following diphtheria-tetanus-pertussis immunization. *Pediatrics* 1988; **81**: 345–9.
5. Shields WD, *et al.* Relationship of pertussis immunization to the onset of neurologic disorders: a retrospective epidemiologic study. *J Pediatr* 1988; **113**: 801–5.
6. Griffin MR, *et al.* Risk of seizures and encephalopathy after immunization with the diphtheria-tetanus-pertussis vaccine. *JAMA* 1990; **263**: 1641–5.
7. Gale JL, *et al.* Risk of serious acute neurological illness after immunization with diphtheria-tetanus-pertussis vaccine: a population-based case-control study. *JAMA* 1994; **271**: 37–41.
8. Miller DL, *et al.* Pertussis immunisation and serious acute neurological illness in children. *BMJ* 1981; **282**: 1595–9.
9. Miller D, *et al.* Pertussis immunisation and serious acute neurological illnesses in children. *BMJ* 1993; **307**: 1171–6.
10. Bellman MH, *et al.* Infantile spasms and pertussis immunisation. *Lancet* 1983; **i**: 1031–4.
11. Howson CP, Fineberg HV. Adverse events following pertussis and rubella vaccines: summary of a report of the Institute of Medicine. *JAMA* 1992; **267**: 392–6.
12. CDC. Update: vaccine side effects, adverse reactions, contraindications, and precautions: recommendations of the Advisory Committee on Immunization Practices (ACIP). *MMWR* 1996; **45** (RR-12): 1–45. Also available at: <http://www.cdc.gov/mmwr/PDF/rr/rr4512.pdf> (accessed 25/05/06)
13. Committee on Infectious Disease, American Academy of Pediatrics. The relationship between pertussis vaccine and central nervous system sequelae: continuing assessment. *Pediatrics* 1996; **97**: 279–81.

Precautions

As for vaccines in general, p.2202. The precautions and contra-indications to the use of pertussis vaccines have sometimes been more stringent than is now considered necessary because of the controversy about their potential adverse effects, especially neurotoxicity (see under Adverse Effects, above). In the UK it is now recommended that immunisation should continue with acellular pertussis vaccine even when episodes of fever (irrespective of severity), hypotonic-hyporesponsive episodes, persistent crying or screaming, or severe local reactions (irrespective of extent) have occurred after a preceding dose. Children who have had a local or general reaction to a whole-cell pertussis vaccine should complete their immunisation with acellular pertussis vaccine.

Whether or not children with a personal or family history of convulsions or epilepsy or who have suffered cerebral damage in the neonatal period should receive pertussis vaccines appears to have been the most difficult question to resolve in the past. In the UK it is now considered that a family history of seizures is not a contra-indication to immunisation. When a child has a history of seizures associated with fever, but no evidence of neurological deterioration, immunisation should proceed as normal; advice on the prevention of fever should be given at the time of immunisation (see Fever and Hyperthermia, p.10 for comments on the prevention of fever after immunisation). Similarly when there is a history of seizures not associated with fever, but no evidence of neurological deterioration, immunisation should proceed as normal. If a seizure associated with fever occurs within 72 hours of immunisation, further immunisation should be deferred until the condition is stable if no underlying cause is found and the child has not recovered completely within 24 hours. Immunisation should also be carried out in children with a history of cerebral damage in the neonatal period unless there is evidence of an evolving neurological abnormality. In children with a neurological problem that is still evolving it is recommended that immunisation should be deferred until the condition is stable. If a child develops encephalopathy or encephalitis within 7 days of immunisation, further immunisation should be deferred until the condition is stable if no underlying cause is found and the child has not recovered completely within 7 days.