

Preparations

Ph. Eur.: Diphtheria Vaccine (Adsorbed); Diphtheria Vaccine (Adsorbed, Reduced Antigen Content).

Proprietary Preparations (details are given in Part 3)

Cz.: Aldiana†; **NZ:** Di Anatoxal.

Diphtheria and Tetanus Vaccines

Vacunas de la difteria y el tétanos.

ATC — J07AM51.

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

Ph. Eur. 6.2 (Diphtheria and Tetanus Vaccine (Adsorbed); Vaccinum Diphtheriae et Tetani Adsorbatum). A preparation of diphtheria formol toxoid and tetanus formol toxoid adsorbed on a mineral carrier. The mineral carrier may be hydrated aluminium phosphate or aluminium hydroxide and the resulting mixture is approximately isotonic with blood. The antigenic properties are adversely affected by certain antimicrobial preservatives particularly those of the phenolic type. It contains not less than 30 international units of diphtheria toxoid and not less than 40 international units of tetanus toxoid 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 DT/Vac/Ads(Child) may be used on the label.

The BP 2008 gives Adsorbed Diphtheria-Tetanus Prophylactic as an approved synonym.

Ph. Eur. 6.2 (Diphtheria and Tetanus Vaccine (Adsorbed, Reduced Antigen(s) Content); Vaccinum Diphtheriae et Tetani Antigeni-o(-is) Minutum). It is diphtheria and tetanus vaccine (adsorbed) containing not less than 2 units of diphtheria toxoid and not less than 20 units of tetanus toxoid per dose.

The BP 2008 states that for a vaccine for use in the UK, the amount of diphtheria toxoid used is adjusted so that the final vaccine contains not more than 2.0 flocculation equivalents of diphtheria toxoid per dose.

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

USP 31 (Diphtheria and Tetanus Toxoids Adsorbed). A sterile suspension prepared by mixing suitable quantities of plain or adsorbed diphtheria toxoid and plain or adsorbed tetanus toxoid and, if plain toxoids are used, an aluminium adsorbing agent. The antigenicity or potency and the proportions of the toxoids are such as to provide an immunising dose of each toxoid in the labelled dose. It should be stored at 2° to 8° and not be allowed to freeze.

USP 31 (Tetanus and Diphtheria Toxoids Adsorbed for Adult Use). A sterile suspension prepared by mixing suitable quantities of adsorbed diphtheria toxoid and adsorbed tetanus toxoid using the same precipitating or adsorbing agent for both toxoids. The antigenicity or potency and the proportions of the toxoids are such as to provide, in the labelled dose, an immunising dose of adsorbed tetanus toxoid and one-tenth of the immunising dose of adsorbed diphtheria toxoid specified for children and not more than 2 Lf of diphtheria toxoid. It should be stored at 2° to 8° and not be allowed to freeze.

Adverse Effects and Precautions

As for vaccines in general, p.2201. See also under Diphtheria Vaccines, above, and Tetanus Vaccines, p.2240. Diphtheria and tetanus vaccines are reported to produce fewer adverse effects than diphtheria, tetanus, and pertussis vaccines (see under Incidence of Adverse Effects, p.2210).

Dose-related effects. A high incidence of adverse effects was reported in teenagers inadvertently given a high-dose diphtheria and tetanus vaccine intended for use in infants.¹ Most reactions were classified as mild or moderately severe, but severe local or systemic reactions occurred in a third of those reporting reactions.

1. Sidebotham PD, Lenton SW. Incidence of adverse reactions after administration of high dose diphtheria with tetanus vaccine to school leavers: retrospective questionnaire study. *BMJ* 1996; **313**: 533-4.

Effects on the nervous system. Encephalopathy more commonly follows vaccination with diphtheria, tetanus, and pertussis vaccine than with diphtheria and tetanus vaccine (p.2210). Several cases of encephalopathy occurred in a small region in Italy in children after immunisation against diphtheria and tetanus,¹ although it was not possible to infer a causal relationship. A case of polyradiculoneuritis has been reported in a patient after the use of a diphtheria-tetanus vaccine and was considered most likely to have been due to the tetanus component.²

1. Greco D. Case-control study on encephalopathy associated with diphtheria-tetanus immunization in Campania, Italy. *Bull WHO* 1985; **63**: 919-25.
2. Holliday PL, Bauer RB. Polyradiculoneuritis secondary to immunization with tetanus and diphtheria toxoids. *Arch Neurol* 1983; **40**: 56-7.

GUILLAIN-BARRÉ SYNDROME. Evidence mainly from case reports and uncontrolled studies favoured a causal relationship between vaccination with diphtheria and tetanus vaccines or single-antigen tetanus vaccines and Guillain-Barré syndrome. The data came primarily from immunocompromised patients.¹ However, a later analysis of active surveillance epidemiological studies of Guillain-Barré syndrome and tetanus

vaccination history concluded that if an association exists, it must be extremely rare and not of public health significance.²

1. Stratton KR, *et al.* Adverse events associated with childhood vaccines other than pertussis and rubella: summary of a report from the Institute of Medicine. *JAMA* 1994; **271**: 1602-5.
2. Tuttle J, *et al.* The risk of Guillain-Barré syndrome after tetanus-toxoid-containing vaccines in adults and children in the United States. *Am J Public Health* 1997; **87**: 2045-8.

Interactions

As for vaccines in general, p.2202.

Uses and Administration

Combined adsorbed diphtheria and tetanus vaccines may be used for active immunisation, although vaccines used for primary immunisation usually combine diphtheria, tetanus, and pertussis, and sometimes also *Haemophilus influenzae* and poliomyelitis. Diphtheria and tetanus vaccines are used in some countries for reinforcing doses after primary immunisation; in the USA they are given to adults every 10 years. For discussion of immunisation schedules, see under Vaccines, p.2202.

The non-adsorbed combined diphtheria and tetanus vaccines have weaker immunogenic properties than adsorbed vaccines and are no longer recommended.

Booster doses. In many countries, booster doses of combined diphtheria and tetanus vaccines are recommended every 10 years, and studies have been conducted to assess whether this is necessary. Since the incidence of clinical diphtheria in many countries in western Europe and North America approaches zero, it had been considered that there was no need for booster doses in adults, despite low antibody titres, so long as the policy of immunisation during infancy was maintained.^{1,2} However, after a report³ of an outbreak of clinical diphtheria in Sweden after a period of many years during which no indigenous cases of diphtheria had occurred and the disease was regarded as being eliminated from the country, the question of immunity in adults and the need for re-immunisation again arose. In the USA, it was considered⁴ that re-immunisation every 10 years with a diphtheria and tetanus combined vaccine was mandatory and that this combined vaccine should be used whenever a tetanus vaccine was indicated as in treating emergency wounds. This policy is also adopted in the UK. Outbreaks of diphtheria in Russia and neighbouring countries⁵ have prompted recommendations for booster doses in travellers to these countries.

1. Mathias RG, Schechter MT. Booster immunisation for diphtheria and tetanus: no evidence of need in adults. *Lancet* 1985; **i**: 1089-91.
2. Anonymous. Diphtheria and tetanus boosters. *Lancet* 1985; **i**: 1081-2.
3. Rappuoli R, *et al.* Molecular epidemiology of the 1984-1986 outbreak of diphtheria in Sweden. *N Engl J Med* 1988; **318**: 12-14.
4. Karzon DT, Edwards KM. Diphtheria outbreaks in immunized populations. *N Engl J Med* 1988; **318**: 41-3.
5. Anonymous. Diphtheria immunisation—advice from the Chief Medical Officer. *Commun Dis Rep* 1993; **3**: 27.

Preparations

Ph. Eur.: Diphtheria and Tetanus Vaccine (Adsorbed); Diphtheria and Tetanus Vaccine (Adsorbed, Reduced Antigen(s) Content);

USP 31: Diphtheria and Tetanus Toxoids Adsorbed; Tetanus and Diphtheria Toxoids Adsorbed for Adult Use.

Proprietary Preparations (details are given in Part 3)

Arg.: Diftavax†; DT Vax†; Imovax DT; Vacuna Doble; **Austral.:** ADT; CDT; **Austria:** DT-reduct; Td-pur; **Belg.:** Ditemer†; Tedivax; **Braz.:** Dif-Tet-All†; DT Vax†; Refortrix†; **Canad.:** Td Adsorbed; **Cz.:** Alditean†; **Denm.:** DiTe Booster; **Fin.:** DiTe Booster; **Ger.:** DT-Impfstoff†; Td-Impfstoff†; Td-pur; Td-Rix; **Gr.:** Anatoxal Di Te Bema†; DT Vax; **Hong Kong:** DiTe Anatoxal†; **India:** DT-Vac Dual Antigen; **Irl.:** Diftavax; **Ital.:** Anatoxal Adult†; Dif-Tet-All†; Diftavax; Ditanrix; **Malaysia:** Di Te Anatoxal†; **Norw.:** DiTe Booster; **NZ:** ADT; CDT; DiTe Anatoxal; **Philipp.:** Di Te Anatoxal; **Pol.:** DT; **S.Afr.:** DT Vax; **Singapore:** Di Te Anatoxal†; **Spain:** Anatoxal Di Te†; Anatoxal Te Di; Diftavax; Ditanrix; **TD.:** DiTe Booster; **Switz.:** Anatoxal Di Te; Ditanrix; **Thai.:** Adsorbed DT Vaccine; Di Te Anatoxal†; Dif-Tet-All†; DT Vax†; **Turk.:** Di Te Anatoxal; **UK:** Diftavax†; **USA:** Decavac.

Diphtheria, Tetanus, and Haemophilus Influenzae Vaccines

Vacunas de la difteria, el tétanos y *Haemophilus influenzae*.

Profile

Combined adsorbed diphtheria, tetanus, and *Haemophilus influenzae* type b vaccines have been used in some countries for active immunisation of infants. For discussion of immunisation schedules see under Vaccines, p.2202. For concern over the antigenicity of *Haemophilus influenzae* type b vaccine in combined vaccines, see under *Haemophilus Influenzae* Vaccines, Interactions, p.2213.

Diphtheria, Tetanus, and Hepatitis B Vaccines

ATC — J07CA07.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus, and Hepatitis B (rDNA) Vac-

cine (Adsorbed); Vaccinum Diphtheriae, Tetani et Hepatitidis B (ADNr) Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tetanus formol toxoid, hepatitis B surface antigen, and 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.

Profile

Combined diphtheria, tetanus, and hepatitis B vaccines have been used in some countries for active immunisation.

Preparations

Ph. Eur.: Diphtheria, Tetanus and Hepatitis B (rDNA) Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Gr.: Primavax†.

Diphtheria, Tetanus, and Pertussis Vaccines

Vacunas de la difteria, el tétanos y la tos ferina.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus and Pertussis Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani et Pertussis Adsorbatum). A preparation of diphtheria formol toxoid and tetanus formol toxoid on a mineral carrier to which a suspension of killed *Bordetella pertussis* has been added. The mineral carrier may be hydrated aluminium phosphate or aluminium hydroxide and the resulting mixture is approximately isotonic with blood. The antigenic properties are adversely affected by certain antimicrobial preservatives particularly those of the phenolic type. It contains not less than 30 international units of diphtheria toxoid, not less than 40 international units if the test is performed in *guinea-pigs*, or 60 international units if the test is performed in *mice*, of tetanus toxoid, and not less than 4 international units of the pertussis component 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 DTWP may be used on the label.

Ph. Eur. 6.2 (Diphtheria, Tetanus and Pertussis (Acellular, Component) Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani et Pertussis Sine Cellulis ex Elementis Praeparatum Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, and 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 BP2008 states that DTaP may be used on the label.

Adverse Effects and Precautions

As for vaccines in general, p.2201. See also under Diphtheria Vaccines, p.2209, Pertussis Vaccines, p.2230, and Tetanus Vaccines, p.2240.

The incidence of local reactions and fever is reported to be lower with the current accelerated immunisation schedules than the formerly used schedules spreading primary immunisation over 6 months. Local reactions and pyrexia occur less commonly after acellular pertussis vaccines than whole-cell pertussis vaccines, especially in children older than 6 months.

In infants with a personal or close family history of seizures, precautions should be taken to avoid pyrexia. See under Pertussis Vaccines for further details of precautions and contra-indications in individuals with a history of neurological problems.

Incidence of adverse effects. The incidence of local reactions is lower with diphtheria and tetanus vaccines combined with an acellular pertussis component (acellular DTP) than with a whole-cell pertussis component, and is similar to that after diphtheria and tetanus (DT) vaccines. Such reactions are generally mild and self-limiting. Rarely, high fever, persistent or inconsolable crying (possibly as a reaction to pain), hypotonic-hyporesponsive collapse, or short-lived convulsions (frequently febrile convulsions) may occur, and have been reported after both DT and acellular DTP vaccines with equal frequency. These reactions do not appear to have any long-term consequences. Rare but serious acute neurological complications including encephalopathy and prolonged seizures have been reported after DTP vaccines and have been attributed to the whole-cell pertussis component (see Effects on the Nervous System, p.2230) but the association could be coincidental. Epidemiological studies have shown that such events are exceedingly rare and only occasionally followed by long-term neurological damage. Analysis of these studies has been difficult but authorities in the UK and USA concluded that the evidence was insufficient for a link.

A causal relationship between DTP vaccination and sudden infant death syndrome (SIDS) has not been established and any temporal relationship is likely to be due to chance.^{1,2} There is evidence that the risk of SIDS is lower in infants who have been vaccinated.³

Immediate anaphylactic reactions have been reported and are regarded as a contra-indication to further use of DTP vaccine. However, the appearance of a rash is not generally regarded as a contra-indication to further doses.

For further details concerning the contra-indications and precautions to be observed for pertussis-containing vaccines, see p.2230.

- Hoffman HJ, et al. Diphtheria-tetanus-pertussis immunization and sudden infant death: results of the National Institute of Child Health and Human Development Cooperative Epidemiological Study of Sudden Infant Death Syndrome Risk Factors. *Pediatrics* 1987; **79**: 598-611.
- Griffin MR, et al. Risk of sudden infant death syndrome after immunisation with the diphtheria-tetanus-pertussis vaccine. *N Engl J Med* 1988; **319**: 618-23.
- Mitchell EA, et al. Immunisation and the sudden infant death syndrome. *Arch Dis Child* 1995; **73**: 498-501.

Interactions

As for vaccines in general, p.2202.

For a report of a diminished immune response to *Haemophilus influenzae* conjugated vaccine when mixed with diphtheria, tetanus, and acellular pertussis vaccine, see Haemophilus Influenzae Vaccines, p.2213.

Uses and Administration

Combined diphtheria, tetanus, and pertussis vaccines are used for active immunisation of children. For discussion of immunisation schedules, see under Vaccines, p.2202.

Combined adsorbed vaccines may be given by deep subcutaneous or intramuscular injection (vaccines with acellular pertussis components are for intramuscular injection only) in usual doses of 0.5 mL. In the USA, a vaccine with an acellular pertussis component is used as part of the recommended schedule for primary immunisation. Three doses are given at intervals of 2 months (the first preferably at 2 months of age), a fourth dose at least 6 months after the third, and a fifth dose at school entry. Another dose of a vaccine specially formulated for use in adults and adolescents is given at 11 to 12 years of age.

Vaccines containing an acellular pertussis component are now preferred to those containing a whole-cell component (see Vaccine Development, p.2231). The non-adsorbed type of combined diphtheria, tetanus, and pertussis vaccines have weaker immunogenic properties than adsorbed vaccines and are no longer recommended.

Preparations

Ph. Eur.: Diphtheria, Tetanus and Pertussis (Acellular, Component) Vaccine (Adsorbed); Diphtheria, Tetanus and Pertussis Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Arg.: Bustrix; Triacel; Vacuna **Bulg.**: Boostrix; Infanrix; Tripacel; **Austria**: Boostrix; Infanrix; **Belg.**: Boostrix; Infanrix; Triamer; **Braz.**: DT-Coq/DTP; Infanrix; Pertacel; Vacina Accl Ads Contra Dif. Tet e Coq; Vacina Comb. Contra Dif-Tet-Pert. Accl Vacina Comb. Contra Dif-Tet-Coq; **Canad.**: Adacel; **Cz.**: Alditeperax; Boostrix; Infanrix; **Denm.**: Di-Te-Ki-Booster; **Fin.**: Boostrix; Di-Te-Kik; Infanrix; **Ger.**: Boostrix; Covaxis; Infanrix; **Gr.**: Anatoxal Di Te Per; DiTePer Anatoxal; DT Coq; Infanrix; **Hong Kong**: Adsorbed DT Coq; Infanrix; Tripacel; Triple Antigen; **India**: Triple Antigen; Tripacel; **Indon.**: Infanrix; Tripacel; **Ir.**: Infanrix; **Israe.**: Accluvax DTP; Boostrix; DT Coq/DTP; Infanrix; **Ital.**: Boostrix; Infanrix; **Malaysia**: Boostrix; Infanrix; Tripacel; **Mex.**: Boostrix; **Neth.**: Infanrix; **Norw.**: Boostrix; **NZ**: Boostrix; DiTePer Anatoxal; Infanrix; Tripacel; Triple Antigen; **Philipp.**: DFT; DT Coq; Infanrix; Tripacel; **Pol.**: DT Coq; DTP; Infanrix DTPa; Tripacel; **Port.**: Boostrix; Infanrix; **Rus.**: Infanrix (Индфаринкс); **S.Afr.**: DTP-Merieux; Infanrix; DTPa; **Singapore**: Boostrix; Infanrix; Tripacel; **Spain**: Anatoxal Di Te Per; Boostrix; DTP-Merieux; Infanrix; **Swed.**: Di-Te-Kik; Infanrix; **Switz.**: Boostrix; Infanrix DTPa; **Thai.**: Boostrix; DT Coq/DTP; DTP Vaccine; Infanrix; Tripacel; **Turk.**: Accluvax DTP; Di Te Per Anatoxal; Infanrix DTPa; Tripacel; **UK**: Infanrix; Trivax-AD; **USA**: Adacel; Boostrix; Daptacel; Infanrix; Tripedia.

Diphtheria, Tetanus, Pertussis, and Haemophilus Influenzae Vaccines

Vacunas de la difteria, el tétanos, la tos ferina y Haemophilus influenzae.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis (Acellular, Component) and Haemophilus type b Conjugate Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum et Haemophilii Stirpe b Conjugatum Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, polyribosylribitol phosphate derived from a suitable strain of *Haemophilus influenzae* type b and covalently bound to a carrier protein, and a mineral carrier such as aluminium hydroxide or hydrated aluminium phosphate. The product may be presented with the Haemophilus type b component in a separate container, the contents of which are mixed with the other components immediately before use. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light.

Adverse Effects and Precautions

As for vaccines in general, p.2201.

See also under Diphtheria Vaccines, p.2209, Diphtheria, Tetanus, and Pertussis Vaccines, p.2210, Haemophilus Influenzae Vac-

cines, p.2213, Pertussis Vaccines, p.2230, and Tetanus Vaccines, p.2240.

Interactions

As for vaccines in general, p.2202.

Uses and Administration

Combined adsorbed diphtheria, tetanus, whole-cell or acellular pertussis, and *Haemophilus influenzae* type b vaccines are available in some countries for active immunisation of children. For discussion of immunisation schedules, see under Vaccines, p.2202. Some combined vaccines are not licensed for use in primary immunisation regimens because of concerns over the response to the *Haemophilus influenzae* type b component (see under Interactions of Haemophilus Influenzae Vaccines, p.2213).

Preparations

Ph. Eur.: Diphtheria, Tetanus, Pertussis (Acellular, Component) and Haemophilus Type b Conjugate Vaccine (Adsorbed); Diphtheria, Tetanus, Pertussis (Acellular, Component), Poliomyelitis (Inactivated) and Haemophilus Type b Conjugate Vaccine (Adsorbed); Diphtheria, Tetanus, Pertussis, Poliomyelitis (Inactivated) and Haemophilus Type b Conjugate Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Arg.: Actacel; **Austral.**: Infanrix Hib; **Belg.**: Infanrix + Hib; **Braz.**: Tetract-Hib; Vacina Comb. Contra Dif-Tet-Pert. Accl e Hib; **Chile**: Actacel; Tetract-Hib; **Cz.**: Infanrix Hib; Tetract-Hib; **Ger.**: Infanrix + Hib; **Indon.**: Infanrix Hib; Tetract-Hib; **Israe.**: Infanrix Hib; Tetract-Hib; **Malaysia**: Infanrix Hib; Tetract-Hib; **NZ**: Infanrix Hib; **Philipp.**: Tetract-Hib; **Port.**: Infanrix Hib; **S.Afr.**: Actacel; Combact-Hib; **Singapore**: Actacel; Infanrix Hib; Tetract-Hib; **Spain**: Infanrix Hib; Tetract-Hib; **Switz.**: Infanrix DTPa-Hib; **Thai.**: Actacel; Tetract-Hib; **Turk.**: Tetract-Hib; **UK**: Act-Hib DTP; Infanrix Hib; **USA**: TriHibit; **Venez.**: Vacuna Tetract-Hib;

Diphtheria, Tetanus, Pertussis, Haemophilus Influenzae, and Hepatitis B Vaccines

ATC — J07CA11.

Profile

Combined diphtheria, tetanus, pertussis, Haemophilus influenzae, and hepatitis B vaccines are available in some countries for active immunisation.

Preparations

Proprietary Preparations (details are given in Part 3)

Arg.: Tritanrix HB-Hib; **Cz.**: Quintanrix; **Mex.**: Tritanrix HB + Hiberix; **Neth.**: Quintanrix; **Port.**: Quintanrix.

Multi-ingredient: **NZ**: Tritanrix HB + Hib.

Diphtheria, Tetanus, Pertussis, and Hepatitis B Vaccines

Vacunas de la difteria, el tétanos, la tos ferina y la hepatitis B.

ATC — J07CA05.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis (Acellular, Component) and Hepatitis B (rDNA) Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum et Hepatitis B (ADNr) Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, hepatitis B surface antigen, and 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.

Profile

Combined diphtheria, tetanus, pertussis, and hepatitis B vaccines are available in some countries for active immunisation.

Preparations

Ph. Eur.: Diphtheria, Tetanus, Pertussis (Acellular, Component) and Hepatitis B (rDNA) Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Austral.: Infanrix HepB; **Braz.**: Vacina Comb. Contra Dif-Tet-Pert-Accl e Hepat.B; **Cz.**: Infanrix HepB; Tritanrix HEPB; **Fin.**: Infanrix HepB; **Gr.**: Infanrix HepB; Infanrix-HepB; **India**: Tritanrix HB; **Indon.**: Tritanrix HB; **Ital.**: Infanrix HepB; **Malaysia**: Tritanrix HB; **Mex.**: Tritanrix HB; **Neth.**: Tritanrix-HepB; **NZ**: Infanrix HepB; **Philipp.**: Tritanrix HB; **S.Afr.**: Infanrix HB; Tritanrix HB; **Spain**: Infanrix HepB; Tritanrix HB; **Swed.**: Infanrix HepB; **Thai.**: Tritanrix HB; **Turk.**: Tritanrix HB; **Venez.**: Tritanrix HB;

Diphtheria, Tetanus, Pertussis, Hepatitis B, Poliomyelitis, and Haemophilus Influenzae Vaccines

ATC — J07CA09.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis (Acellular, Component), Hepatitis B (rDNA), Poliomyelitis (Inactivated) and Haemophilus type b Conjugate Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum, Hepatitis B (ADNr), Poliomyelitis Inactivatum et Haemophilii Stirpe b Coniugatum Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, hepatitis B surface antigen, suitable strains of human polioviruses type 1, 2, and 3 grown in suitable cell cultures and inactivated by a validated method, polyribosylribitol phosphate derived from a suitable strain of *Haemophilus influenzae* type b and co-

valently bound to a carrier protein, and a mineral carrier such as aluminium hydroxide or hydrated aluminium phosphate. The product may be presented with the Haemophilus type b component in a separate container, the contents of which are mixed with the other components immediately before or during use. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light.

Profile

A combined diphtheria, tetanus, pertussis, hepatitis B, poliomyelitis, and Haemophilus influenzae vaccine is available in some countries for active immunisation.

References

- Curran MP, Goa KL. DTPa-HBV-IPV/Hib vaccine (Infanrix hexa). *Drugs* 2003; **63**: 673-82.

Preparations

Ph. Eur.: Diphtheria, Tetanus, Pertussis (Acellular, Component), Hepatitis B (rDNA), Poliomyelitis (Inactivated) and Haemophilus Type b Conjugate Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Arg.: Hexavac; Infanrix Hexa; **Austral.**: Infanrix Hexa; **Austria**: Hexavac; Infanrix Hexa; **Belg.**: Infanrix Hexa; **Braz.**: Hexavac; Infanrix Hexa; Vacina Adsorvida Contra Dif, Tet, Coq Accl, Polio Inat Hepat B (Rec) e Hib Coni; Vacina Comb. Contra Dif-Tet-Pert. Accl Hepat.B r-DNA, Polio Inat e Hib; **Chile**: Hexavac; Infanrix Hexa; **Cz.**: Infanrix Hexa; **Fin.**: Infanrix Hexa; **Fr.**: Hexavac; Infanrixhexa; **Ger.**: Hexavac; Infanrix Hexa; **Gr.**: Hexavac; Infanrix Hexa; Infanrix-HepB/HI; **Hong Kong**: Infanrix Hexa; **Hung.**: Hexavac; **Ital.**: Hexavac; Infanrix Hexa; **Malaysia**: Infanrix Hexa; **Mex.**: Hexavac; Infanrix Hexa; **Neth.**: Infanrix Hexa; **NZ**: Infanrix Hexa; **Philipp.**: Infanrix Hexa; **Pol.**: Hexavac; Infanrix Hexa; **Port.**: Infanrix Hexa; **Singapore**: Infanrix Hexa; **Spain**: Hexavac; Infanrix Hexa; **Swed.**: Hexavac; Infanrix Hexa; **Switz.**: Hexavac; Infanrix Hexa; **Thai.**: Infanrix Hexa; **Venez.**: Infanrix Hexa.

Diphtheria, Tetanus, Pertussis, and Poliomyelitis Vaccines

Vacunas de la difteria, el tétanos, la tos ferina y la poliomieltis.

ATC — J07CA02.

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

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis (Acellular, Component) and Poliomyelitis (Inactivated) Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum et Poliomyelitis Inactivatum Adsorbatum). A combined vaccine containing diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, suitable strains of human polioviruses type 1, 2, and 3 grown in suitable cell cultures and inactivated by a validated method, and 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.

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis and Poliomyelitis (Inactivated) Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis et Poliomyelitis Inactivatum Adsorbatum). A combined vaccine containing diphtheria formol toxoid, tetanus formol toxoid, an inactivated suspension of *Bordetella pertussis*, suitable strains of human polioviruses type 1, 2, and 3 grown in suitable cell cultures and inactivated by a validated method, and 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.

Ph. Eur. 6.2 (Diphtheria, Tetanus, Pertussis (Acellular, Component) and Poliomyelitis (Inactivated) Vaccine (Adsorbed, Reduced Antigen(s) Content); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum et Poliomyelitis Inactivatum, Antigeni- α (-is) Minutum, Adsorbatum). A combined vaccine containing diphtheria formol toxoid, tetanus formol toxoid, individually purified antigenic components of *Bordetella pertussis*, suitable strains of human polioviruses type 1, 2, and 3 grown in suitable cell cultures and inactivated by a validated method, and a mineral adsorbent such as aluminium hydroxide or hydrated aluminium phosphate. The amount of diphtheria toxoid per single human dose is reduced compared to vaccines generally used for primary vaccination; the amounts of tetanus toxoid and pertussis components may also be reduced. It should be stored at 2° to 8°, not be allowed to freeze, and be protected from light.

Adverse Effects and Precautions

As for vaccines in general, p.2201.

See also under Diphtheria Vaccines, p.2209, Diphtheria, Tetanus, and Pertussis Vaccines, p.2210, Pertussis Vaccines, p.2230, and Tetanus Vaccines, p.2240.

Interactions

As for vaccines in general, p.2202.

Uses and Administration

A combined diphtheria, tetanus, pertussis (acellular component), and poliomyelitis (inactivated) vaccine is used for active immunisation. For discussion of immunisation schedules see under Vaccines, p.2202.

In the UK it is used as part of the recommended schedule and is given by intramuscular injection in a single dose of 0.5 mL as a booster at pre-school age (3 years