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 and studgen infant death: results of the National Institute of Child Health and Human Development Cooperative Epidemiological Study of Sudden Infant Death Syndrome Risk Factors. *Pediat-*rics 1987; **79:** 598–611.
- Griffin MR, et al. Risk of sudden infant death syndrome after immunization 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 Triple; Austral.: Boostrix, Infanrix, Tripacel;
Austria: Boostrix, Infanrix; Belg.: Boostrix, Infanrix; Triamer; Braz.: DTCoq/DTP; Infanrix; Pertacel; Vacina Acel Ads Contra Dif. Tet e Coq.
Vacina Comb. Contra Dif.-Tet.-Pert. Acel; Vacina Comb. Contra Dift.-Tet. Vacina Comb. Contra Dif. Tet., Pert. Acel; Vacina Comb. Contra Dift. Tet.
Coq. Acel; Canad: Adacel; Cz.: Alditeperaț; Boostrix, Infanrix; Denns:
Di-Te.-Ki-Boostre; Fin.: Boostrix; Di-Te.-Kiţh; Infanrix; Ger.: Boostrix; Covaxis; Infanrix; Gr.: Anatoxal Di Te Perţ; Di-TePer Anatoxalţ; DT Coqţ; Infanrix; Hong Kong: Adsorbed DT Coqţ; Infanrix; Tipacel; Tiple Antigenţ; Infanrix; Tipacel; Tiple Antigenţ; Infanrix; Tipacel; Infanrix; Infanrix; Rorw:: Boostrix; Infanrix; DT-Coqţ; Di-Te, Infanrix; Infanrix; Norw:: Boostrix; NT: Boostrix; DT-Pere Anatoxal; Infanrix; Tipacel; Tiple
Antigen; Philipps.: DPT, DT COq; Infanrix; Tipacel; Pol.: DT Coq; DI-P.
DTP; Infanrix DTPa; Tripacel; Port.: Boostrix; Infanrix; Infanrix; Infanrix; Infanrix; Swed.: Di-Te-Kik; Infanrix; Swet.: Di-Te-Kik; Infanrix; Swet.: Di-Te-Kik; Infanrix; Swet.: Di-Te-Kik; Infanrix; DTPa; Tipacel; Tiple: Boostrix; DTPA; Tipacel; Tiple: Boostrix; DTPA; Tipacel; Tiple: Boostrix; DTPA; Tipacel; Infanrix; Tipacel; Tiple: Acelluxax DTP; Di Te Per Anatoxal; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tirvax-AD; USA: Adacel; Boostrix; Daptacel; Infanrix; Tipacel; UK: Infanrix; Tipacel; UK: Infanrix; Tipacel; UK: Infanrix;

# Diphtheria, Tetanus, Pertussis, and **Haemophilus Influenzae Vaccines**

Vacunas de la difteria, el tétanos, la tos ferina y Haemophilus in-

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 Haemophili Stirpe b Conjugatum Adsorbatum). A combined vaccine composed of diphtheria formol toxoid, tet-anus 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 Vaccines, 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.: Actace!; Austral.: Infanrix Hib; Belg.: Infanrix + Hib†; Braz.: Tetract-HiB†; Vacina Comb. Contra Dif. Tet.-Pert. Acel e HiB; Chile: Actace!; Tetract-HiB; Yacina Comb. Contra Dif. Tet.-Pert. Acel e HiB; Chile: Actace!; Tetract-HiB; Deta: Infanrix Hib; Tetract-HiB; Acel: Infanrix Hib; Tetract-HiB; Malaysia: Infanrix Hib; Tetract-HiB; Acel: Infanrix Hib; Tetract-HiB; Acel: Infanrix Hib; Tetract-HiB; Spain: Infanrix Hib; Tetract-HiB; Spain: Infanrix Hib; Tetract-HiB; Tetract

# 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

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 Hepatitidis 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

#### **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.Acel e HepatB; Cz.: Infanrix HepB; Tritanrix HEPB; Fin.: Infanrix HepB; Gritanrix HepB; Fin.: Infanrix HepB; Gritanrix HepB; Fin.: Infanrix HepB; Gritanrix HB; Infanrix HepB; Infanrix Hep; Infanrix HepB; Infanrix HepB; Malaysia: Tritanrix HB; Mex.: Tritanrix HB; Neth.: Tritanrix-HepB; NZ: Infanrix HepB; Philipp.: Tritanrix HB; Svefi: Infanrix HepB; Tritanrix HB; Svefi: Infanrix HB; Tritanrix HB; Trit

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

ATC - 107CA09.

Pharmacopoeias. Many pharmacopoeias, including Eur. (see p.vii), have monographs. **Ph. Eur. 6.2** ( Diphtheria, Tetanus, Pertussis (Acellular, Compo-

nent), Hepatitis B (rDNA), Poliomyelitis (Inactivated) and Haemohenti, hepatius B (rDNA), Polioniyellis (inactivated) and Haerito-philus type b Conjugate Vaccine (Adsorbed); Vaccinum Diphthe-riae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum, Hepatitidis B (ADNr), Poliomyelitidis Inactivatum et Haemophili Stirpe b Conjugatum 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 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 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†; Arg.: Hexavac†, Infanrix Hexa; Austral: Infanrix Hexa; Austral: Hexavac†, Infanrix Hexa; Belg.: Infanrix Hexa; Braz.: Hexavac†, Infanrix Hexa; Vacina Adsorvida Contra Dif, Tet, Coq Acel, Polio Inat Hepat B (Rec) e Hib Conj; Vacina Comb. Contra Dif. Tet, Pert. Acel Hepat B r-DNA, Polio Inat e Hibs; Callie: Hexavac†, Infanrix Hexa; Fr.: Hexavac†, InfanrixHexa; Ger.: Hexavac†, Infanrix Hexa; Gr.: Hexavac†, InfanrixHexa; Ger.: Hexavac†, Infanrix Hexa; Gr.: Hexavac†, Infanrix Hexa; Molaysia: Infanrix Hexa; Hexavac†, Infanrix Hexa; Molaysia: Infanrix Hexa; Pol.: Hexavac†, Infanrix Hexa; Swed.: Hexavac†, Infanrix Hexavac†, In

# Diphtheria, Tetanus, Pertussis, and Poliomyelitis Vaccines

Vacunas de la difteria, el tétanos, la tos ferina y la poliomielitis. ATC — J07CA02.

Pharmacopoeias. Many pharmacopoeias, including Eur. (see p.vii), have monographs. **Ph. Eur. 6.2** (Diphtheria, Tetanus, Pertussis (Acellular, Compo-

nent) and Poliomyelitis (Inactivated) Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum et Poliomyelitidis 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, Tétani, Pertussis et Poliomyelitidis 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 Poliomyelitidis Inactivatum, Antigeni-o(-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 per-tussis 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 4 months to 5 years). It is not suitable for primary immunisation.

#### **Preparations**

**Ph. Eur.:** Diphtheria, Tetanus, Pertussis (Acellular, Component) and Poliomyelitis (Inactivated) Vaccine (Adsorbed); Diphtheria, Tetanus, Pertussis (Acellular, Component) and Poliomyelitis (Inactivated) Vaccine (Adsorbed, Reduced Antigen(s) Content); Diphtheria, Tetanus, Pertussis and Poliomy elitis (Inactivated) Vaccine (Adsorbed).

Proprietary Preparations (details are given in Part 3)

Propriecary Preparations (details are given in Part 3)

Austral: Bosotrix (PV; Infanrix (PV; Quadrace; Austria: Repevax; Tetravac;
Belg: Infanrix (PV; Tetracoq†; Tetravac; Braz.: Tetracoq†; Vacina Acel Ads
Contra Dif, Tet, Coq e Polio Inat Comb CVac Conj Contra Hib; Canad.:
Quadrace); Cz.: Infanrix Polio; Denm.: Di-Te-Ki-Pol; Fin.: Boostrix Polio; Quadracel; Cz.: Infanrix Polic; Denm.: Di-Te-Ki-Pol; Fin.: Boostrix Polic; Di-Te-Ki-Pol; Infanrix Polic; Tetravac; Fiz. Boostrixtera: Infanrixetar: Repevax; Tetravac; Ger.: Boostrix Polic; Quatro-Virelon†; Repevax; Tetravac†; Gr.: Boostrix Polic; Infanrix Tetra Repevax; Tetravac; Hall: Tetravac; Infanrix IPV; Tetracq†; Tetraxim; Mex.: Infanrix IPV; Neth.: Infanrix IPV; Triaxis; Norw.: Boostrix Polic; NZ: Boostrix IPV; Infanrix IPV; Quadracel; Philipp.: Tetracq; Tetraxim; Pol.: DTaP-IPV; Tetracq†; Port.: Boostrix Polic; Infanrix IPV; Tetravac; Tetravac; Swet.: Boostrix Polic; Di-Ki-Pol†; Tetravac; Swet.: Boostrix Polic; Di-Ki-Pol†; Tetravac; Switz.: Boostrix Polic; Infanrix IPV; Repevax; Venez.: Vacuna Adsorbida Tetravalente. sorbida Tetravalente

# Diphtheria, Tetanus, Pertussis, Poliomyelitis, and Haemophilus Influenzae Vaccines

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

- J07CA06.

Pharmacopoeias. Many pharmacopoeias, including Eur. (see

p.vii), have monographs. **Ph. Eur. 6.2** (Diphtheria, Tetanus, Pertussis (Acellular, Component), Poliomyelitis (Inactivated) and Haemophilus type b Conjugate Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis Sine Cellulis ex Elementis Praeparatum Poliomyelitidis Inactivatum et Haemophili Stirpe b Conjugatum Adsorbatum). A combined vaccine composed of 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, 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 hy-droxide or hydrated aluminium phosphate. The product is 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.

**Ph. Eur. 6.2** (Diphtheria, Tetanus, Pertussis, Poliomyelitis (Inactivated) and Haemophilus type b Conjugate Vaccine (Adsorbed); Vaccinum Diphtheriae, Tetani, Pertussis, Poliomyelitidis Inactivatum et Haemophili Stirpe b Conjugatum Adsorbatum). A combined vaccine composed of 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, polyri-bosylribitol 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 is 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 Vaccines, p.2213, Pertussis Vaccines, p.2230, and Tetanus Vaccines, p.2240.

Premature neonates. In an observational study<sup>1</sup> of 78 verylow-birth-weight premature neonates given a combined diphtheria, tetanus, pertussis (acellular component), poliomyelitis (inactivated), and Haemophilus influenzae vaccine before hospital discharge, increased incidences of apnoea, bradycardia, desaturation, or oxygen requirement occurred in 47% overall within 24 to 48 hours of vaccination. All neonates with increased events returned to baseline within 48 to 72 hours and there was no detrimental impact on clinical course. The authors considered that, although monitoring and appropriate intervention were required delaying vaccination was not warranted, a view in line with UK and USA official recommendations (see p.2202).

1. Pfister RE, et al. Safety of DTaP-based combined immunization in very-low-birth-weight premature infants: frequent but mostly benign cardiorespiratory events. *J Pediatr* 2004; **145:** 58–66.

#### Interactions

As for vaccines in general, p.2202.

#### **Uses and Administration**

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

In the UK it is used as part of the recommended schedule for primary immunisation. It is given by intramuscular injection in usual doses of 0.5 mL; three doses are given at intervals of one month, starting preferably at 2 months of age. Although it is not licensed for use after a child's fourth birthday, the national schedule considers it may be used up to the age of 10 years.

#### **Preparations**

Proprietary Preparations (details are given in Part 3)

Arg.: Pentasim; Poliacel†; Austral.: Pediacel; Poliacel; Austria: Infanrix IPV + Hib; Belg: Infanrix IPV + Hib; Braz: Infanrix IPV + Hib; Pentact-HIB; Poliacel†; Vacina Comb. Contra Dif.-Tet.-Pert. Acel, Polio Inat e HIB; Canda: Pentacel: Chile: Pentact-HIB; Canda: Pentacel: Chile: Pentact-HIB; Ca.: Infanrix IPV + Hib; Denm.: Di-Tenad.: Pentacel: Chile: Pentact-HIB; Cz.: Infanrix IPV + Hib; Denm.: Di-Te-Ki-Pol/Act-Hib; Fin.: Infanrix Polio + Hib; Pentavac; Fr.: Infanrix quinta; Pentavacq; Gr.: Infanrix IPV + Hib; Pentavac; Gr.: Infanrix IPV + Hib; Pentavac; Gr.: Infanrix IPV + Hib; Pentavac; Hong Kong: Infanrix IPV + Hib; Pentavac; Hong Kong: Infanrix IPV + Hib; Pentavac; Pentavim; III: Infanrix IPV + Hib; Pentavac; Pentavim; III: Infanrix IPV + Hib; Pentavim; Mex.: Infanrix IPV + Hib; Pediace; Neth.: DKT-Hib; Infanrix IPV + Hib; Pediace; Neth.: DKT-Hib; Infanrix IPV + Hib; Pediace; Neth.: DKT-Bib; Pentavim; IPV + Hib; Pentavim; IPV + Hib; Pentavim; IPV + Hib; Pentavim; IIIV + Hib; Pentavim; IIIII: Infanrix IPV + Hib; Pentavim; IIII: Infanrix IPV + Hib; Vacuna Adoptida Pentavalente. Venez.: Infanrix IPV + Hib; Vacuna Adsorbida Pentavalente

### Diphtheria, Tetanus, Pertussis, Poliomyelitis, and Hepatitis B Vaccines

ATC - J07CA12

#### **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, Hepatitis B Vaccines, p.2215, 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), poliomyelitis (inactivated), and hepatitis B vaccine is available in some countries for active immunisation of children.

# **Preparations**

Proprietary Preparations (details are given in Part 3) Austral.: Infanrix Penta; Cz.: Infanrix Penta; Gr.: Infanrix Penta; Ital.: Infanrix Penta; Neth.: Infanrix Penta; USA: Pediarix

# Diphtheria, Tetanus, and **Poliomyelitis Vaccines**

Vacunas de la difteria, el tétanos y la poliomielitis. ATC — J07CA01.

Pharmacopoeias. Many pharmacopoeias, including Eur. (see

p.vii), have monographs. **Ph. Eur. 6.2** ( Diphtheria, Tetanus and Poliomyelitis (Inactivated) Vaccine (Adsorbed, Reduced Antigens(s) Content); Vaccinum Diphtheriae, Tetani et Poliomyelitidis Inactivatum, Antigeni-o(-is) Minutum, Adsorbatum). A combined vaccine containing diph theria formol toxoid, tetanus formol toxoid, suitable strains of human polioviruses types 1, 2, and 3 grown in suitable cell cul-tures 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 amount of tetanus toxoid 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 and Tetanus Vaccines, p.2210, and Tetanus Vaccines, p.2240.

#### Interactions

As for vaccines in general, p.2202.

# **Uses and Administration**

A combined diphtheria, tetanus, 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 (usually 0.5 mL) as a booster at the ages of 13 to 18 years. It is not licensed for primary immunisation.

#### **Preparations**

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

Proprietary Preparations (details are given in Part 3)

Austria: Revaxis; Belg.: Revaxis; Canad.: Td-Polio; Fr.: DT Polio; Revaxis;
Vaccin DTP†; Ger.: Revaxis; Td-Virelon; Gr.: Revaxis; Hung.: Dultavax; Irl.:
Revaxis; Ital.: Revaxis, Weth.: Revaxis; Port.: Revaxis; Switz.: Revaxis; Td-Virelon: **UK:** Revaxis.

#### **Endotoxin Antibodies**

Anticuerpos antiendotoxinas

#### Profile

Antibodies against the endotoxin of Gram-negative bacteria have been tried as adjunctive therapy for the treatment and prevention of Gram-negative bacteraemia and shock.

Early preparations consisted of antisera prepared from the sera of donors immunised with *Escherichia coli* J5; these were superseded by human and murine IgM monoclonal antibodies. Nebacumab (HA-1A) is a human monoclonal IgM antibody that binds specifically to the lipid A domain of endotoxin. Lipid A in the circulation releases tumour necrosis factor and other cytokines from macrophages and endothelial cells which may ultimately culminate in physiological effects such as multiple organ failure. Despite early promising results of clinical studies the safety of nebacumab in patients without Gram-negative septicae-mia was questioned and the product was withdrawn.

A murine monoclonal IgM antibody (edobacomab; E5) has also undergone clinical trials although results have been disappoint-

### **Epstein-Barr Virus Vaccines**

Vacunas del virus de Epstein-Barr.

#### **Profile**

Several Epstein-Barr virus vaccines are under investigation for active immunisation against infectious mononucleosis and posttransplant lymphoproliferative disorders.

♦ Epstein-Barr virus is a herpesvirus that is ubiquitous in the adult population. It only causes clinical illness where primary infection occurs in adolescence or adulthood, when it prompts the symptoms of infectious mononucleosis in about 50% of cases. More than 90% of the world's population, however, carry the virus as a lifelong latent infection of B-lymphocytes and, as a result, Epstein-Barr virus can also be associated with malignancies including lymphoproliferative diseases, Burkitt's lymphoma, gastric carcinoma, oral hairy leucoplakia, nasopharyngeal carcinoma, and Hodgkin's disease

Vaccines against Epstein-Barr virus infection are under investigation<sup>1,2</sup> and the main focus has been towards the development of a vaccine to prevent primary infection or to minimise its consequences, namely infectious mononucleosis and posttransplant lymphoproliferative disease, rather than towards the malignancies associated with the virus which occur in relatively fewer patients. Two main approaches have been adopted, the first of which seeks to exploit the major envelope glycoprotein of the virus, gp340, because of its ability to induce neutralising antibodies. This vaccine may prevent infectious mononucleosis by moderating the initial viral replication and spread during primary infection, thereby curtailing the cytotoxic T-lymphocyte response to lytic antigens that would otherwise invoke the immunological processes responsible for clinical symptoms. The second approach is based on the induction of cytotoxic T-cells specific to Epstein-Barr virus, thereby aiming to reduce the clinical symptoms of infectious mononucleosis rather than to prevent primary infection.

Potential future vaccines for malignancies associated with Epstein-Barr virus are likely to be therapeutic rather than preventative and to exploit the presence of the virus in tumour cells; alternatively they may be focussed on tumour antigens not encoded by Epstein-Barr virus. <sup>1,2</sup>

- Moss DJ, et al. Candidate vaccines for Epstein-Barr virus. BMJ 1998; 317: 423-4.
   Macsween KF, Crawford DH. Epstein-Barr virus—recent advances. Lancet Infect Dis 2003; 3: 131-40.

# **Escherichia Coli Vaccines**

Vacunas de Escherichia coli.

Vaccines against enterotoxigenic strains of Escherichia coli are under investigation. Vaccine candidates include toxoids, inactivated whole bacteria, purified surface antigens, and live oral vac-

 $\Diamond$  Infectious diarrhoea remains a major source of morbidity and mortality in the world and a significant proportion is caused by pathogenic strains of *Escherichia coli* While it is considered feasible to develop effective vaccines against *E. coli*, at present there are no such vaccines available. Current approaches against enter-opathogenic *E. coli* (EPEC) and enterohaemorrhagic *E. coli* (EHEC) have focussed on three main areas: the EPEC and EHEC proteins involved in colonisation of the intestine, the EHEC O157-specific side-chain of lipopolysaccharides, and the