

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use HIBERIX safely and effectively. See full prescribing information for HIBERIX.

HIBERIX [Haemophilus b Conjugate Vaccine (Tetanus Toxoid Conjugate)]

Solution for Intramuscular Injection
Initial U.S. Approval: 2009

RECENT MAJOR CHANGES

Indications and Usage (1)	xx/xxxx
Dosage and Administration, Reconstitution (2.1)	xx/xxxx
Dosage and Administration, Dose and Schedule (2.3)	xx/xxxx
Warnings and Precautions, Apnea in Premature Infants (5.3)	xx/xxxx

INDICATIONS AND USAGE

HIBERIX is a vaccine indicated for active immunization for the prevention of invasive disease caused by *Haemophilus influenzae* type b. HIBERIX is approved for use in children 6 weeks through 4 years of age (prior to fifth birthday). (1)

No clinical data are available from controlled studies comparing booster immunization with HIBERIX and a US-licensed Haemophilus b Conjugate Vaccine. (1)

DOSAGE AND ADMINISTRATION

A 4-dose series (0.5 mL each) given by intramuscular injection (2.3):

- Primary series: One dose each at 2, 4, and 6 months of age. The first dose may be given as early as 6 weeks of age.
- Booster: One dose at 15 through 18 months of age.

DOSAGE FORMS AND STRENGTHS

Solution for injection supplied as a vial of lyophilized vaccine to be reconstituted with the accompanying vial of saline diluent. A single dose, after reconstitution, is 0.5 mL. (3)

CONTRAINDICATIONS

Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any *H. influenzae* type b- or tetanus toxoid-containing vaccine or any component of HIBERIX. (4)

WARNINGS AND PRECAUTIONS

- If Guillain-Barré syndrome has occurred within 6 weeks of receipt of a prior vaccine containing tetanus toxoid, the decision to give HIBERIX should be based on potential benefits and risks. (5.1)
- Syncope (fainting) can occur in association with administration of injectable vaccines, including HIBERIX. Procedures should be in place to avoid falling injury and to restore cerebral perfusion following syncope. (5.2)
- Apnea following intramuscular vaccination has been observed in some infants born prematurely. Decisions about when to administer an intramuscular vaccine, including HIBERIX, to infants born prematurely should be based on consideration of the individual infant's medical status, and the potential benefits and possible risks of vaccination. (5.3)

ADVERSE REACTIONS

Common solicited adverse events ($\geq 20\%$) were pain and redness at the injection site, irritability, drowsiness, fever, loss of appetite, fussiness, and restlessness. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact GlaxoSmithKline at 1-888-825-5249 or VAERS at 1-800-822-7967 or www.vaers.hhs.gov.

DRUG INTERACTIONS

Do not mix HIBERIX with any other vaccine in the same syringe or vial. (7.2)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: XX/XXXX

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1 FULL PRESCRIBING INFORMATION

2 1 INDICATIONS AND USAGE

3 HIBERIX[®] is indicated for active immunization for the prevention of invasive disease caused by
4 *Haemophilus influenzae* type b. HIBERIX is approved for use in children 6 weeks through 4
5 years of age (prior to fifth birthday).

6 The evaluation of effectiveness of HIBERIX was based on immune responses in children using
7 serological endpoints that predict protection from invasive disease due to *H. influenzae* type b
8 [see *Clinical Pharmacology (12.1)*, *Clinical Studies (14.1)*]. These protective antibody levels
9 have not been evaluated in clinical trials in which a booster dose of HIBERIX is compared to a
10 booster dose of a US-licensed Haemophilus b Conjugate Vaccine in children who previously
11 received a primary series with a US-licensed Haemophilus b Conjugate Vaccine [see *Clinical*
12 *Studies (14.1)*].

13 2 DOSAGE AND ADMINISTRATION

14 2.1 Reconstitution

15 HIBERIX is to be reconstituted only with the accompanying saline diluent. The reconstituted
16 vaccine should be a clear and colorless solution. Parenteral drug products should be inspected
17 visually for particulate matter and discoloration prior to administration, whenever solution and
18 container permit. If either of these conditions exists, the vaccine should not be administered.



Figure 1. Cleanse both vial stoppers. Withdraw 0.6 mL of saline diluent from accompanying vial.



Figure 2. Transfer 0.6 mL saline diluent into lyophilized vaccine vial.



Figure 3. Shake the vial well.



Figure 4. After reconstitution, withdraw 0.5 mL of reconstituted vaccine and administer **intramuscularly**.

19 Use a separate sterile needle and sterile syringe for each individual.

20 After reconstitution, administer HIBERIX immediately or store refrigerated between 2° and 8°C
21 (36° and 46°F) and administer within 24 hours. If the vaccine is not administered immediately,
22 shake the solution well again before administration.

23 **2.2 Administration**

24 **For intramuscular use only.**

25 HIBERIX is administered as a single dose (0.5 mL) by intramuscular injection into the
26 anterolateral aspect of the thigh or deltoid.

27 Do not administer this product intravenously, intradermally, or subcutaneously.

28 **2.3 Dose and Schedule**

29 HIBERIX is administered as a 4-dose series (0.5-mL each dose) given by intramuscular
30 injection. The series consists of a primary immunization course of 3 doses administered at 2, 4,
31 and 6 months of age, followed by a booster dose administered at 15 through 18 months of age.
32 The first dose may be given as early as 6 weeks of age.

33 **3 DOSAGE FORMS AND STRENGTHS**

34 HIBERIX is a solution for injection supplied as a single-dose vial of lyophilized vaccine to be
35 reconstituted with the accompanying vial of saline diluent. A single dose, after reconstitution, is
36 0.5 mL.

37 **4 CONTRAINDICATIONS**

38 Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any *H. influenzae* type b- or
39 tetanus toxoid-containing vaccine or any component of the vaccine is a contraindication to
40 administration of HIBERIX [see *Description (11)*].

41 **5 WARNINGS AND PRECAUTIONS**

42 **5.1 Guillain-Barré Syndrome**

43 If Guillain-Barré syndrome has occurred within 6 weeks of receipt of a prior vaccine containing
44 tetanus toxoid, the decision to give any tetanus toxoid-containing vaccine, including HIBERIX,
45 should be based on careful consideration of the potential benefits and possible risks.

46 **5.2 Syncope**

47 Syncope (fainting) can occur in association with administration of injectable vaccines, including
48 HIBERIX. Syncope can be accompanied by transient neurological signs such as visual
49 disturbance, paresthesia, and tonic-clonic limb movements. Procedures should be in place to
50 avoid falling injury and to restore cerebral perfusion following syncope.

51 **5.3 Apnea in Premature Infants**

52 Apnea following intramuscular vaccination has been observed in some infants born prematurely.
53 Decisions about when to administer an intramuscular vaccine, including HIBERIX, to infants
54 born prematurely should be based on consideration of the individual infant's medical status, and
55 the potential benefits and possible risks of vaccination.

56 **5.4 Preventing and Managing Allergic Vaccine Reactions**

57 Prior to administration, the healthcare provider should review the patient's immunization history
58 for possible vaccine hypersensitivity. Epinephrine and other appropriate agents used for the
59 control of immediate allergic reactions must be immediately available should an acute
60 anaphylactic reaction occur.

61 **5.5 Altered Immunocompetence**

62 Safety and effectiveness of HIBERIX in immunosuppressed children have not been evaluated. If
63 HIBERIX is administered to immunosuppressed children, including children receiving
64 immunosuppressive therapy, the expected immune response may not be obtained.

65 **5.6 Interference with Laboratory Tests**

66 Urine antigen detection may not have a diagnostic value in suspected disease due to
67 *H. influenzae* type b within 1 to 2 weeks after receipt of a *H. influenzae* type b-containing
68 vaccine, including HIBERIX [see *Drug Interactions (7.1)*].

69 **5.7 Tetanus Immunization**

70 Immunization with HIBERIX does not substitute for routine tetanus immunization.

71 **6 ADVERSE REACTIONS**

72 **6.1 Clinical Trials Experience**

73 Because clinical trials are conducted under widely varying conditions, adverse reaction rates
74 observed in the clinical trials of a vaccine cannot be directly compared with rates in the clinical
75 trials of another vaccine, and may not reflect the rates observed in practice. There is the
76 possibility that broad use of HIBERIX could reveal adverse reactions not observed in clinical
77 trials.

78 Across clinical trials, common solicited adverse events ($\geq 20\%$) were pain and redness at the
79 injection site, irritability, drowsiness, fever, loss of appetite, fussiness, and restlessness.

80 Study 1: In a randomized, controlled clinical trial conducted in the US, children were vaccinated
81 with HIBERIX (N = 2,963), a US-licensed monovalent Haemophilus b Conjugate Vaccine
82 (Control PRP-T) (Sanofi Pasteur SA) (N = 520), or a US-licensed combined Diphtheria and
83 Tetanus Toxoids and Acellular Pertussis Adsorbed, Inactivated Poliovirus and Haemophilus b
84 Conjugate Vaccine (DTaP-IPV/Hib) (Sanofi Pasteur Ltd.) (N = 520) at 2, 4, and 6 months of age.
85 HIBERIX and Control PRP-T (Sanofi Pasteur SA) were administered concomitantly with
86 PEDIARIX[®] (DTaP-HBV-IPV) [Diphtheria and Tetanus Toxoids and Acellular Pertussis
87 Adsorbed, Hepatitis B (Recombinant) and Inactivated Poliovirus Vaccine] and Pneumococcal
88 13-valent Conjugate Vaccine (PCV13) (Wyeth Pharmaceuticals Inc.) with Doses 1, 2, and 3 and
89 ROTARIX[®] [Rotavirus Vaccine, Live, Oral] with Doses 1 and 2. DTaP-IPV/Hib was
90 administered concomitantly with PCV13 and ENGERIX-B[®] [Hepatitis B Vaccine

91 (Recombinant)] with Doses 1, 2, and 3 and ROTARIX with Doses 1 and 2. If a birth dose of
92 hepatitis B vaccine was received, ENGERIX-B was given with Doses 1 and 3. In the total
93 population, 51.2% were male; 61% were white, 8% were Asian, 9% were black, and 22% were
94 other racial/ethnic groups.

95 In 7 additional clinical studies, 1,008 children received HIBERIX as a booster dose following
96 primary vaccination with either HIBERIX (N = 530), Haemophilus b Conjugate Vaccine
97 (Control PRP-T) (Sanofi Pasteur SA) (N = 235), Haemophilus b Conjugate Vaccine (Merck &
98 Co., Inc.) (N = 26), or Haemophilus b Conjugate Vaccine (Wyeth Pharmaceuticals Inc.) (no
99 longer licensed in the US, N = 217). None of the studies included a comparator group that
100 received a booster dose with a US-licensed Haemophilus b Conjugate Vaccine. Studies were
101 conducted in Europe, Canada, and Latin America. Across these studies, the mean age of subjects
102 at the time of booster vaccination with HIBERIX ranged from 16 to 19 months. At the time of
103 vaccination, 172 (17.1%) subjects were 11 to 14 months of age, 642 (63.7%) subjects were 15 to
104 18 months of age, and 194 (19.2%) subjects were 19 to 25 months of age. Approximately half of
105 the subjects were male. Among subjects for whom information on race/ethnicity was available,
106 nearly all subjects were white.

107 In these 7 studies, HIBERIX was administered concomitantly with non-US formulations
108 (containing 2.5 mg 2-phenoxyethanol per dose as preservative) of one of the following US-
109 licensed vaccines: INFANRIX[®] (DTaP) [Diphtheria and Tetanus Toxoids and Acellular Pertussis
110 Vaccine Adsorbed], KINRIX[®] (DTaP-IPV) [Diphtheria and Tetanus Toxoids and Acellular
111 Pertussis Adsorbed and Inactivated Poliovirus Vaccine], or PEDIARIX (DTaP-HBV-IPV). In the
112 studies, DTaP-IPV and DTaP-HBV-IPV were administered in dosing regimens not approved in
113 the US. Some subjects received DTaP-HBV (GlaxoSmithKline Biologicals, not licensed in US)
114 concomitantly with HIBERIX.

115 Solicited Adverse Events

116 The reported frequencies of solicited local and general adverse events from Study 1 are presented
117 in Table 1.

118 **Table 1. Percentage of Children with Solicited Local and General Adverse Events within**
 119 **4 Days of Primary Series Vaccination^a (at 2, 4, and 6 Months of Age) with HIBERIX^b,**
 120 **Control PRP-T^b, or DTaP-IPV/Hib^c, Total Vaccinated Cohort^d**

Adverse Events	HIBERIX			Control PRP-T			DTaP-IPV/Hib		
	%			%			%		
	Dose			Dose			Dose		
	1	2	3	1	2	3	1	2	3
Local^e									
N	2,828	2,668	2,553	498	481	463	492	469	443
Pain	49.4	45.1	42.8	57.2	53.2	48.2	58.1	50.1	48.5
Pain, grade 3 ^f	3.9	2.7	1.9	9.0	5.4	3.5	8.9	3.2	2.7
Redness	18.7	25.4	29.4	23.5	32.0	29.6	25.6	30.7	37.0
Redness, >20 mm	0.9	0.7	0.7	2.2	1.0	0.2	2.0	2.1	2.3
Swelling	13.0	15.4	18.7	18.5	21.8	19.7	19.5	23.7	23.7
Swelling, >20 mm	1.5	1.0	0.8	4.2	2.7	0.6	3.9	1.9	2.0
General									
N	2,830	2,669	2,553	499	480	463	492	469	443
Irritability	68.9	70.4	67.1	76.4	71.0	67.2	73.0	66.7	69.3
Irritability, grade 3 ^g	4.1	6.4	4.8	8.4	7.7	5.2	6.1	4.5	3.2
Drowsiness	59.9	54.1	49.3	65.7	55.6	49.5	60.6	51.8	49.7
Drowsiness, grade 3 ^h	2.4	2.8	2.2	3.8	2.1	1.3	3.9	2.6	2.7
Loss of appetite	28.7	28.3	27.6	33.3	31.5	27.2	33.5	24.3	24.2
Loss of appetite, grade 3 ⁱ	0.7	1.6	1.5	2.0	1.0	0.4	0.6	0.4	0.5
Fever	13.7	19.2	18.7	16.4	18.8	16.2	11.6	10.9	17.8
Fever, grade 3 ^j	0.3	0.6	0.7	0.4	0.4	0.9	0.0	0.0	0.5

121 N = All subjects for whom safety data were available.

122 ^a Within 4 days of vaccination defined as day of vaccination and the next 3 days.

123 ^b Each dose (Doses 1, 2, and 3) of HIBERIX or Control PRP-T (Sanofi Pasteur SA) was
 124 concomitantly administered with PEDIARIX (DTaP-HBV-IPV) and PCV13. Doses 1 and 2 were
 125 concomitantly administered with ROTARIX.

126 ^c Each dose (Doses 1, 2, and 3) of DTaP-IPV/Hib was concomitantly administered with PCV13 and
 127 ENGERIX-B with Doses 1, 2, and 3 and ROTARIX with Doses 1 and 2. If a birth dose of hepatitis B
 128 vaccine was received, ENGERIX-B was given with Doses 1 and 3.

129 ^d Study 1: NCT01000974.

130 ^e Local reactions at the injection site for HIBERIX, Control PRP-T, or DTaP-IPV/Hib.

131 ^f Grade 3 pain defined as cried when limb was moved/spontaneously painful.

132 ^g Grade 3 irritability defined as crying that could not be comforted/prevented normal activity.

133 ^h Grade 3 drowsiness defined as prevented normal daily activity.

134 ⁱ Grade 3 loss of appetite defined as did not eat at all.

135 ^j Fever defined as $\geq 100.4^{\circ}\text{F}$ ($\geq 38.0^{\circ}\text{C}$) rectally; Grade 3 fever defined as $> 103.1^{\circ}\text{F}$ ($> 39.5^{\circ}\text{C}$) rectally.

136 In an open-label, multicenter study conducted in Germany (Study 2), 371 children received a
 137 booster dose of HIBERIX administered concomitantly with DTaP-HBV-IPV. The mean age at
 138 the time of vaccination was 16 months. Subjects in this study had previously received a primary
 139 series with either HIBERIX (N = 92), Control PRP-T (Sanofi Pasteur SA) (N = 96), or
 140 Haemophilus b Conjugate Vaccine (Wyeth Pharmaceuticals Inc.) (no longer licensed in the US)
 141 (N = 183). All subjects previously received 3 doses of DTaP-HBV-IPV. The reported
 142 frequencies of solicited local and general adverse events are presented in Table 2.

143 **Table 2. Percentage of Children with Solicited Local and General Adverse Events**
 144 **within 4 Days of Booster Vaccination^a (Dose 4) with HIBERIX^b Coadministered**
 145 **with DTaP-HBV-IPV^c, Intent-to-Treat Cohort (N = 371)**

Adverse Events	% Any	% Grade 3
Local^d		
Redness	24.5	2.4 ^e
Pain	20.5	1.1 ^f
Swelling	14.8	2.2 ^e
General		
Fever ^g	34.8	3.8
Fussiness	25.9	0.8 ^h
Loss of appetite	22.9	0.8 ⁱ
Restlessness	21.8	0.5 ⁱ
Sleepiness	19.9	1.1 ⁱ
Diarrhea	14.6	0.8 ⁱ
Vomiting	4.9	0.5 ⁱ

146 N = All subjects for whom safety data were available.

147 ^a Within 4 days of vaccination defined as day of vaccination and the next 3 days.

148 ^b In this study, 92 subjects previously received 3 doses of HIBERIX, 96 subjects previously
 149 received 3 doses of a Control PRP-T (Sanofi Pasteur SA), and 183 subjects previously
 150 received 3 doses of a Haemophilus b Conjugate Vaccine that is no longer licensed in the US.

151 ^c In this study, DTaP-HBV-IPV was given to subjects who previously received 3 doses of
 152 DTaP-HBV-IPV. In the US, PEDIARIX is approved for use as a 3-dose primary series; use as
 153 a fourth consecutive dose is not approved in the US.

154 ^d Local reactions at the injection site for HIBERIX.

155 ^e Grade 3 redness or swelling defined as >20 mm.

156 ^f Grade 3 pain defined as causing crying when limb moved.

157 ^g Fever defined as $\geq 100.4^{\circ}\text{F}$ ($\geq 38.0^{\circ}\text{C}$) rectally or $\geq 99.5^{\circ}\text{F}$ ($\geq 37.5^{\circ}\text{C}$) axillary, oral, or
 158 tympanic; Grade 3 fever defined as $> 103.1^{\circ}\text{F}$ ($> 39.5^{\circ}\text{C}$) rectally or $> 102.2^{\circ}\text{F}$ ($> 39.0^{\circ}\text{C}$)
 159 axillary, oral, or tympanic.

160 ^h Grade 3 fussiness defined as persistent crying and could not be comforted.

161 ⁱ Grade 3 for these symptoms defined as preventing normal daily activity.

162 Serious Adverse Events

163 In Study 1, one of 2,963 subjects who received HIBERIX and coadministered vaccines given at
164 2, 4, and 6 months of age experienced a SAE which was in temporal association with vaccination
165 and had no alternative plausible causes (convulsion on Day 14 after Dose 1).

166 In the 7 additional studies, two of 1,008 subjects reported a serious adverse event that occurred in
167 the 31-day period following booster immunization with HIBERIX. One subject developed
168 bilateral pneumonia 9 days post-vaccination and one subject experienced asthenia following
169 accidental drug ingestion 18 days post-vaccination.

170 **6.2 Postmarketing Experience**

171 In addition to reports in clinical trials, worldwide voluntary reports of adverse events received
172 for HIBERIX since market introduction (1996) of this vaccine are listed below. This list includes
173 serious events and/or events which have a plausible causal connection to HIBERIX. Because
174 these events are reported voluntarily from a population of uncertain size, it is not possible to
175 reliably estimate their frequency or establish a causal relationship to vaccination.

176 General Disorders and Administration Site Conditions

177 Extensive swelling of the vaccinated limb, injection site induration.

178 Immune System Disorders

179 Allergic reactions (including anaphylactic and anaphylactoid reactions), angioedema.

180 Nervous System Disorders

181 Convulsions (with or without fever), hypotonic-hyporesponsive episode (i.e., sudden onset of
182 hypotonia, hyporesponsiveness, and pallor or cyanosis), somnolence, syncope, or vasovagal
183 responses to injection.

184 Respiratory, Thoracic, and Mediastinal Disorders

185 Apnea [*see Warnings and Precautions (5.3)*].

186 Skin and Subcutaneous Tissue Disorders

187 Rash, urticaria.

188 **7 DRUG INTERACTIONS**

189 **7.1 Interference with Laboratory Tests**

190 Haemophilus b capsular polysaccharide derived from Haemophilus b Conjugate Vaccines has
191 been detected in the urine of some vaccinees.¹ Urine antigen detection may not have a diagnostic
192 value in suspected disease due to *H. influenzae* type b within 1 to 2 weeks after receipt of a

193 *H. influenzae* type b-containing vaccine, including HIBERIX [see *Warnings and Precautions*
194 (5.6)].

195 **7.2 Concomitant Vaccine Administration**

196 In Study 1, HIBERIX was administered concomitantly with PEDIARIX (DTaP-HBV-IPV),
197 PCV13, and ROTARIX [see *Adverse Reactions (6.1)*, *Clinical Studies (14.2)*].

198 In the 7 additional studies, a booster dose of HIBERIX was administered concomitantly with 1 of
199 the following vaccines: DTaP, DTaP-IPV, DTaP-HBV-IPV, or DTaP-HBV (GlaxoSmithKline
200 Biologicals, not licensed in the US). The formulations of DTaP, DTaP-IPV, and DTaP-HBV-IPV
201 were non-US formulations (containing 2.5 mg 2-phenoxyethanol per dose as preservative) of the
202 following US-licensed vaccines: INFANRIX, KINRIX, and PEDIARIX, respectively. In these
203 studies, DTaP-IPV and DTaP-HBV-IPV were administered in dosing regimens that are not
204 approved in the US. [See *Adverse Reactions (6.1)*, *Clinical Studies (14.1)*.]

205 If HIBERIX is administered concomitantly with other injectable vaccines, they should be given
206 with separate syringes and at different injection sites. HIBERIX should not be mixed with any
207 other vaccine in the same syringe or vial.

208 **7.3 Immunosuppressive Therapies**

209 Immunosuppressive therapies, including irradiation, antimetabolites, alkylating agents, cytotoxic
210 drugs, and corticosteroids (used in greater than physiologic doses), may reduce the immune
211 response to HIBERIX.

212 **8 USE IN SPECIFIC POPULATIONS**

213 **8.1 Pregnancy**

214 Pregnancy Category C

215 Animal reproduction studies have not been conducted with HIBERIX. It is also not known
216 whether HIBERIX can cause fetal harm when administered to a pregnant woman or can affect
217 reproduction capacity.

218 **8.4 Pediatric Use**

219 Safety and effectiveness of HIBERIX in children younger than 6 weeks of age and in children 5
220 to 16 years of age have not been established.

221 **11 DESCRIPTION**

222 HIBERIX [Haemophilus b Conjugate Vaccine (Tetanus Toxoid Conjugate)] is a solution for
223 intramuscular injection, supplied as a sterile, lyophilized powder which is reconstituted at the
224 time of use with the accompanying saline diluent. HIBERIX contains Haemophilus b capsular
225 polysaccharide (polyribosyl-ribitol-phosphate [PRP]), a high molecular weight polymer prepared
226 from the *Haemophilus influenzae* type b strain 20,752 grown in a synthetic medium that

227 undergoes heat inactivation and purification. The tetanus toxin, prepared from *Clostridium tetani*
228 grown in a semi-synthetic medium, is detoxified with formaldehyde and purified. The capsular
229 polysaccharide is covalently bound to the tetanus toxoid. After purification, the conjugate is
230 lyophilized in the presence of lactose as a stabilizer. The diluent for HIBERIX is a sterile saline
231 solution (0.9% sodium chloride) supplied in vials.

232 After reconstitution, each 0.5-mL dose is formulated to contain 10 mcg of purified capsular
233 polysaccharide conjugated to approximately 25 mcg of tetanus toxoid, 12.6 mg of lactose, and
234 ≤0.5 mcg of residual formaldehyde.

235 HIBERIX does not contain a preservative.

236 The lyophilized vaccine and saline diluent vial stoppers are not made with natural rubber latex.

237 **12 CLINICAL PHARMACOLOGY**

238 **12.1 Mechanism of Action**

239 *Haemophilus influenzae* is a gram-negative coccobacillus. Most strains of *H. influenzae* that
240 cause invasive disease are type b. *H. influenzae* type b can cause invasive disease such as sepsis
241 and meningitis.

242 Specific levels of antibodies to polyribosyl-ribitol-phosphate (anti-PRP) have been shown to
243 correlate with protection against invasive disease due to *H. influenzae* type b. Based on data from
244 passive antibody studies² and a clinical efficacy study with unconjugated *Haemophilus* b
245 polysaccharide vaccine³, an anti-PRP concentration of 0.15 mcg/mL has been accepted as a
246 minimal protective level. Data from an efficacy study with unconjugated *Haemophilus* b
247 polysaccharide vaccine indicate that an anti-PRP concentration of ≥1.0 mcg/mL predicts
248 protection through at least a 1-year period.^{4,5} These antibody levels have been used to evaluate
249 the effectiveness of *Haemophilus* b Conjugate Vaccines, including HIBERIX.

250 **13 NONCLINICAL TOXICOLOGY**

251 **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

252 HIBERIX has not been evaluated for carcinogenic or mutagenic potential, or for impairment of
253 fertility.

254 **14 CLINICAL STUDIES**

255 **14.1 Immunological Evaluation**

256 Primary Series Vaccination (Doses 1, 2, and 3)

257 The immunogenicity of HIBERIX was evaluated in a randomized, controlled trial (Study 1).
258 HIBERIX or control vaccines were administered concomitantly with US-licensed vaccines [*see*
259 *Adverse Reactions (6.1)*].

260 Anti-PRP GMCs and seroprotection rates 1 month following Dose 3 of HIBERIX, Control PRP-
 261 T (Sanofi Pasteur SA), or DTaP-IPV/Hib are presented in Table 3.

262 **Table 3. Anti-PRP GMCs and Seroprotection Rates 1 Month following 3 Doses of**
 263 **HIBERIX, Control PRP-T^a, or DTaP-IPV/Hib^b Administered at 2, 4, and 6 Months**
 264 **of Age, ATP Cohort for Immunogenicity^c**

Vaccine	N	Anti-PRP GMC (mcg/mL) (95% CI)	% Anti-PRP ≥0.15 mcg/mL (95% CI)	% Anti-PRP ≥1.0 mcg/mL (95% CI)
HIBERIX	1,590	5.19 (4.77, 5.66)	96.6 (95.6, 97.4)	81.2 (79.2, 83.1)
Control PRP-T	274	6.74 (5.59, 8.13)	96.7 ^d (93.9, 98.5)	89.8 ^e (85.6, 93.1)
DTaP-IPV/Hib	253	3.64 (2.89, 4.58)	92.5 ^f (88.5, 95.4)	78.3 ^f (72.7, 83.2)

265 ^a US-licensed monovalent Haemophilus b Conjugate Vaccine (Control PRP-T) (Sanofi Pasteur
 266 SA).

267 ^b US-licensed DTaP-IPV/Hib Vaccine (Sanofi Pasteur Ltd.).

268 ^c Study 1: NCT01000974.

269 ^d HIBERIX was non-inferior to Control PRP-T for percent of subjects achieving anti-PRP
 270 ≥0.15 mcg/mL [lower limit of 95% CI on difference of HIBERIX minus Control PRP-T ≥
 271 predefined limit of -5%].

272 ^e The non-inferiority criterion was not met (lower limit of 95% CI for the difference in the
 273 percentages of subjects with anti-PRP ≥1.0 mcg/mL between two groups [HIBERIX minus
 274 Control PRP-T] was -12.28%, which was lower than the predefined limit of -10%).

275 ^f Analyses of anti-PRP immune responses following DTaP-IPV/Hib vaccination were
 276 exploratory.

277 Booster Vaccination (Dose 4)

278 In 6 clinical studies, the immune response to HIBERIX administered as a booster dose was
 279 evaluated in a total of 415 children 12 to 23 months of age. At the time of vaccination, 30
 280 children were 12 to 14 months of age, 316 children were 15 to 18 months of age, and 69 children
 281 were 19 to 23 months of age. Among subjects, 43% to 60% were male. Among subjects for
 282 whom information on race/ethnicity was available, nearly all subjects were white. None of the
 283 studies included a comparator group that received a booster dose with a US-licensed
 284 Haemophilus b Conjugate Vaccine. Characteristics of 3 of these studies are presented in Table 4.

285 **Table 4. Characteristics of 3 Open-Label Booster Immunization Studies of**
 286 **HIBERIX**

Study	Country	Per-Protocol Immunogenicity Cohort N	Priming History	Booster Vaccination with HIBERIX	
				Age at Vaccination (months)	Concomitantly Administered Vaccine ^a
3	Canada	42	DTaP-HBV-IPV ^b + Haemophilus b Conjugate Vaccine ^c at 2, 4, and 6 months of age	16-18	DTaP-HBV-IPV ^b
4	Canada	64	DTaP-IPV ^d + HIBERIX at 2, 4, and 6 months of age	16-19	DTaP-IPV ^d
5	Germany	108	DTaP-HBV ^e + HIBERIX at 3, 4, and 5 months of age	16-23	DTaP-HBV ^e

287 ^a Administered at a separate site.

288 ^b Non-US formulation equivalent to PEDIARIX with the exception of containing 2.5 mg 2-
 289 phenoxyethanol per dose as preservative. In the US, PEDIARIX is approved for use as a 3-
 290 dose primary series; use as a fourth consecutive dose is not approved in the US.

291 ^c US-licensed Haemophilus b Conjugate Vaccine (Control PRP-T) (Sanofi PasteurSA).

292 ^d Non-US formulation equivalent to KINRIX with the exception of containing 2.5 mg 2-
 293 phenoxyethanol per dose as preservative. In the US, KINRIX is approved for use as the fifth
 294 dose of DTaP and the fourth dose of IPV in children 4 to 6 years of age previously primed
 295 with approved dosing regimens of INFANRIX and/or PEDIARIX. The DTaP-IPV dosing
 296 regimen is not approved in the US.

297 ^e Manufactured by GlaxoSmithKline Biologicals (not licensed in the US).

298 Antibodies to PRP were measured in sera obtained immediately prior to and 1 month after
 299 booster vaccination with HIBERIX. Geometric mean concentrations and anti-PRP seroprotection
 300 rates are presented in Table 5.

301 **Table 5. Anti-PRP GMCs and Seroprotection Rates prior to and 1 Month following**
 302 **a Booster Dose of HIBERIX, Per-Protocol Immunogenicity Cohort**

Study	N	Anti-PRP GMC (mcg/mL)		% Anti-PRP ≥ 0.15 mcg/mL		% Anti-PRP ≥ 1.0 mcg/mL	
		Pre-	Post-	Pre-	Post-	Pre-	Post-
3 ^a	42	0.46	59.07	76.2	100	35.7	97.6
4 ^b	63-64	0.25	47.78	71.4	100	12.7	100
5 ^c	108	0.59	96.12	77.8	100	32.4	100

303 GMC = Geometric mean antibody concentration.

304 N = Number of children for whom serological results were available for the pre- and post-dose
 305 immunological evaluations.

306 Studies 3, 4, and 5 correspond to Studies 3, 4, and 5, respectively in Table 4.

307 ^a Canadian study in children 16 to 18 months of age who previously received 3 doses of DTaP-
 308 HBV-IPV and Haemophilus b Conjugate Vaccine (Control PRP-T) (Sanofi Pasteur SA). The
 309 booster dose of HIBERIX was coadministered with DTaP-HBV-IPV (a fourth consecutive
 310 dose of PEDIARIX is not approved in the US). In this study, pre-vaccination sera may have
 311 been obtained up to 1 week prior to booster vaccination with HIBERIX.

312 ^b Canadian study in children 16 to 19 months of age who previously received 3 doses of DTaP-
 313 IPV and HIBERIX. The booster dose of HIBERIX was coadministered with DTaP-IPV. The
 314 DTaP-IPV dosing regimen is not approved in the US.

315 ^c German study in children 16 to 23 months of age who previously received 3 doses of DTaP-
 316 HBV (GlaxoSmithKline Biologicals, not licensed in the US) and HIBERIX. The booster dose
 317 of HIBERIX was coadministered with DTaP-HBV.

318 **14.2 Concomitant Vaccine Administration**

319 Primary Series Vaccination (Doses 1, 2, and 3)

320 In US Study 1, subjects who received HIBERIX concomitantly with PEDIARIX (DTaP-HBV-
 321 IPV) and PCV13 at 2, 4, and 6 months of age had no evidence for reduced antibody responses
 322 relative to the response in control subjects administered Control PRP-T (Sanofi Pasteur SA)
 323 concomitantly with PEDIARIX (DTaP-HBV-IPV) and PCV13, to pertussis antigens (GMC to
 324 pertussis toxin, filamentous hemagglutinin, and pertactin), diphtheria toxoid (antibody levels
 325 ≥ 0.1 IU/mL), tetanus toxoid (antibody levels ≥ 0.1 IU/mL), poliovirus types 1, 2, and 3 (antibody
 326 levels $\geq 1:8$ to each virus), PCV13 (antibody levels ≥ 0.2 mcg/mL and GMC to each serotype), or
 327 hepatitis B (anti-hepatitis B surface antigen ≥ 10 mIU/mL). The immune responses to PEDIARIX
 328 (DTaP-HBV-IPV) and PCV13 were evaluated 1 month following Dose 3. Subjects in both
 329 groups received ROTARIX at 2 and 4 months of age.

330 **Booster Vaccination (Dose 4)**

331 In 7 additional studies, a booster dose of HIBERIX was administered concomitantly with non-
332 US formulations of INFANRIX, KINRIX, and PEDIARIX. Non-US formulations of KINRIX
333 and PEDIARIX were administered in dosing regimens not approved in the US.

334 Sufficient data are not available to confirm lack of interference in immune responses to vaccines
335 administered concomitantly with a booster dose of HIBERIX.

336 **15 REFERENCES**

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350 **16 HOW SUPPLIED/STORAGE AND HANDLING**

351 HIBERIX is available in single-dose vials of lyophilized vaccine, accompanied by vials
352 containing 0.85 mL of saline diluent (packaged without syringes or needles).

353 Supplied as package of 10 doses (NDC 58160-818-11):

354 NDC 58160-816-01 Vial of lyophilized vaccine in Package of 10: NDC 58160-816-05

355 NDC 58160-817-01 Vial of saline diluent in Package of 10: NDC 58160-817-05

356 **16.1 Storage before Reconstitution**

357 Lyophilized vaccine vials: Store refrigerated between 2° and 8°C (36° and 46°F). Protect vials
358 from light.

359 Diluent: Store refrigerated or at controlled room temperature between 2° and 25°C (36° and
360 77°F). Do not freeze. Discard if the diluent has been frozen.

361 **16.2 Storage after Reconstitution**

362 Administer within 24 hours of reconstitution. After reconstitution, store refrigerated between 2°
363 and 8°C (36° and 46°F). Discard the reconstituted vaccine if not used within 24 hours. Do not
364 freeze. Discard if the vaccine has been frozen.

365 **17 PATIENT COUNSELING INFORMATION**

- 366 • Inform parents or guardians of the potential benefits and risks of immunization with
367 HIBERIX.
- 368 • Inform parents or guardians about the potential for adverse reactions that have been
369 temporally associated with administration of HIBERIX or other vaccines containing similar
370 components.
- 371 • Give parents or guardians the Vaccine Information Statements, which are required by the
372 National Childhood Vaccine Injury Act of 1986 to be given prior to immunization. These
373 materials are available free of charge at the Centers for Disease Control and Prevention
374 (CDC) website (www.cdc.gov/vaccines).

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