

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

KINRIX™ (Diphtheria and Tetanus Toxoids and Acellular Pertussis Adsorbed and Inactivated Poliovirus Vaccine)

Suspension for Intramuscular Injection

Initial U.S. Approval: 2008

INDICATIONS AND USAGE

A single dose of KINRIX is indicated for active immunization against diphtheria, tetanus, pertussis, and poliomyelitis as the fifth dose in the diphtheria, tetanus, and acellular pertussis (DTaP) vaccine series and the fourth dose in the inactivated poliovirus vaccine (IPV) series in children 4 through 6 years of age whose previous DTaP vaccine doses have been with INFANRIX and/or PEDIARIX for the first three doses and INFANRIX for the fourth dose. (1)

DOSAGE AND ADMINISTRATION

A single intramuscular injection (0.5 mL). (2.2)

DOSAGE FORMS AND STRENGTHS

Single-dose vial and prefilled syringe containing a 0.5-mL suspension for injection of diphtheria and tetanus toxoids, acellular pertussis antigens, and inactivated poliovirus types 1, 2, and 3. (3)

CONTRAINDICATIONS

- Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any diphtheria toxoid, tetanus toxoid, pertussis or poliovirus-containing vaccine, or to any component of KINRIX, including neomycin and polymyxin B. (4.1)
- Encephalopathy within 7 days of administration of a previous pertussis-containing vaccine. (4.2)
- Progressive neurologic disorders. (4.3)

WARNINGS AND PRECAUTIONS

- If Guillain-Barré syndrome occurs within 6 weeks of receipt of a prior vaccine containing tetanus toxoid, the decision to give KINRIX should be based on potential benefits and risks. (5.1)
- The needleless prefilled syringes contain dry natural latex rubber and may cause allergic reactions. (5.2)
- If adverse events (i.e., temperature $\geq 105^{\circ}\text{F}$, collapse or shock-like state, persistent, inconsolable crying lasting ≥ 3 hours, occurring within 48 hours of vaccination; seizures within 3 days of vaccination) have occurred in temporal relation to receipt of a pertussis-containing vaccine, the decision to give KINRIX should be based on potential benefits and risks. (5.3)
- For children at higher risk for seizures, an antipyretic may be administered at the time of vaccination with KINRIX. (5.4)

ADVERSE REACTIONS

- The most frequently reported solicited local reaction ($>50\%$) was injection site pain. Other common solicited local reactions ($\geq 25\%$) were redness, increase in arm circumference, and swelling. (6.1)
- Common solicited general adverse events ($\geq 15\%$) were drowsiness, fever ($\geq 99.5^{\circ}\text{F}$), and loss of appetite. (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 KINRIX with any other vaccine in the same syringe or vial. (7.1)

See 17 for PATIENT COUNSELING INFORMATION.

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KNX:2PI

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2 FULL PRESCRIBING INFORMATION

3 1 INDICATIONS AND USAGE

4 A single dose of KINRIX is indicated for active immunization against diphtheria, tetanus,
5 pertussis, and poliomyelitis as the fifth dose in the diphtheria, tetanus, and acellular pertussis
6 (DTaP) vaccine series and the fourth dose in the inactivated poliovirus vaccine (IPV) series in
7 children 4 through 6 years of age whose previous DTaP vaccine doses have been with
8 INFANRIX[®] (Diphtheria and Tetanus Toxoids and Acellular Pertussis Vaccine Adsorbed)
9 and/or PEDIARIX[®] [Diphtheria and Tetanus Toxoids and Acellular Pertussis Adsorbed,
10 Hepatitis B (Recombinant) and Inactivated Poliovirus Vaccine Combined] for the first three
11 doses and INFANRIX for the fourth dose.

12 2 DOSAGE AND ADMINISTRATION

13 2.1 Preparation for Administration

14 Shake vigorously to obtain a homogeneous, turbid, white suspension. Do not use if
15 resuspension does not occur with vigorous shaking. Parenteral drug products should be inspected
16 visually for particulate matter and discoloration prior to administration, whenever solution and
17 container permit. After removal of the dose, any vaccine remaining in the vial should be
18 discarded.

19 2.2 Recommended Dose and Schedule

20 KINRIX is to be administered as a 0.5-mL dose by intramuscular injection. The preferred
21 site of administration is the deltoid muscle of the upper arm. Do not administer this product
22 intravenously, intradermally, or subcutaneously.

23 KINRIX may be used for the fifth dose in the DTaP immunization series and the fourth
24 dose in the IPV immunization series in children 4 through 6 years of age (prior to the seventh
25 birthday) whose previous DTaP vaccine doses have been with INFANRIX and/or PEDIARIX for
26 the first three doses and INFANRIX for the fourth dose [*see Indications and Usage (1)*].

27 3 DOSAGE FORMS AND STRENGTHS

28 KINRIX is available in 0.5-mL single-dose vials and prefilled TIP-LOK[®] syringes.

29 Each 0.5-mL dose contains a suspension for injection of diphtheria and tetanus toxoids,
30 acellular pertussis antigens, and inactivated poliovirus types 1, 2, and 3. See *Description (11)* for
31 the complete listing of ingredients.

32 4 CONTRAINDICATIONS

33 4.1 Hypersensitivity

34 Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any diphtheria toxoid,
35 tetanus toxoid, pertussis or poliovirus-containing vaccine, or to any component of KINRIX,
36 including neomycin and polymyxin B, is a contraindication to administration of KINRIX [*see*

37 *Description (11)*]. Because of the uncertainty as to which component of the vaccine might be
38 responsible, no further vaccination with any of these components should be given. Alternatively,
39 such individuals may be referred to an allergist for evaluation if immunization with any of these
40 components is considered.

41 **4.2 Encephalopathy**

42 Encephalopathy (e.g., coma, decreased level of consciousness, prolonged seizures) within
43 7 days of administration of a previous dose of a pertussis-containing vaccine that is not
44 attributable to another identifiable cause is a contraindication to administration of any pertussis-
45 containing vaccine, including KINRIX.

46 **4.3 Progressive Neurologic Disorder**

47 Progressive neurologic disorder, including infantile spasms, uncontrolled epilepsy, or
48 progressive encephalopathy is a contraindication to administration of any pertussis-containing
49 vaccine, including KINRIX. Pertussis vaccine should not be administered to individuals with
50 such conditions until a treatment regimen has been established and the condition has stabilized.

51 **5 WARNINGS AND PRECAUTIONS**

52 **5.1 Guillain-Barré Syndrome**

53 If Guillain-Barré syndrome occurs within 6 weeks of receipt of a prior vaccine containing
54 tetanus toxoid, the decision to give any tetanus toxoid-containing vaccine, including KINRIX,
55 should be based on careful consideration of the potential benefits and possible risks. When a
56 decision is made to withhold tetanus toxoid, other available vaccines should be given, as
57 indicated.

58 **5.2 Latex**

59 The tip cap and the rubber plunger of the needleless prefilled syringes contain dry natural
60 latex rubber that may cause allergic reactions in latex sensitive individuals. The vial stopper is
61 latex-free.

62 **5.3 Adverse Events Following Prior Pertussis Vaccination**

63 If any of the following events occur in temporal relation to receipt of a pertussis-
64 containing vaccine, the decision to give any pertussis-containing vaccine, including KINRIX,
65 should be based on careful consideration of the potential benefits and possible risks:

- 66 • Temperature of $\geq 40.5^{\circ}\text{C}$ (105°F) within 48 hours not due to another identifiable cause;
- 67 • Collapse or shock-like state (hypotonic-hyporesponsive episode) within 48 hours;
- 68 • Persistent, inconsolable crying lasting ≥ 3 hours, occurring within 48 hours;
- 69 • Seizures with or without fever occurring within 3 days.

70 When a decision is made to withhold pertussis vaccination, other available vaccines
71 should be given, as indicated.

72 **5.4 Children at Risk for Seizures**

73 For children at higher risk for seizures than the general population, an appropriate
74 antipyretic may be administered at the time of vaccination with a pertussis-containing vaccine,
75 including KINRIX, and for the ensuing 24 hours to reduce the possibility of post-vaccination

76 fever.

77 **5.5 Preventing and Managing Allergic Vaccine Reactions**

78 Prior to administration, the healthcare provider should review the patient’s immunization
79 history for possible vaccine sensitivity and previous vaccination-related adverse reactions to
80 allow an assessment of benefits and risks. Epinephrine and other appropriate agents used for the
81 control of immediate allergic reactions must be immediately available should an acute
82 anaphylactic reaction occur.

83 **6 ADVERSE REACTIONS**

84 **6.1 Clinical Trials Experience**

85 Because clinical trials are conducted under widely varying conditions, adverse reaction
86 rates observed in the clinical trials of a vaccine cannot be directly compared with rates in the
87 clinical trials of another vaccine, and may not reflect the rates observed in practice.

88 A total of 3,537 children were vaccinated with a single dose of KINRIX in 3 clinical
89 trials. Of these, 381 children received a non-US formulation of KINRIX (containing ≤ 2.5 mg
90 2-phenoxyethanol per dose as preservative). The primary study (Study 048), conducted in the
91 United States, was a randomized, controlled clinical trial in which children 4 to 6 years of age
92 were vaccinated with KINRIX (N = 3,156) or control vaccines (INFANRIX and IPOL[®] vaccine
93 [IPV, Sanofi Pasteur SA]; N = 1,053) as a fifth DTaP vaccine dose following 4 doses of
94 INFANRIX and as a fourth IPV dose following 3 doses of IPOL. Subjects also received the
95 second dose of US-licensed measles, mumps, and rubella (MMR) vaccine (Merck & Co., Inc.)
96 administered concomitantly, at separate sites.

97 Data on adverse events were collected by parents/guardians using standardized forms for
98 4 consecutive days following vaccination with KINRIX or control vaccines (i.e., day of
99 vaccination and the next 3 days). The reported frequencies of solicited local reactions and
100 general adverse events in Study 048 are presented in Table 1.

101 In 3 studies (Study 046, 047, and 048), children were monitored for unsolicited adverse
102 events, including serious adverse events, that occurred in the 31-day period following
103 vaccination and in 2 studies (Study 047 and 048), parents/guardians were actively queried about
104 changes in the child’s health status, including the occurrence of serious adverse events, through
105 6 months post vaccination.

106

107 **Table 1. Percentage of Children 4 to 6 Years of Age Reporting Solicited Local Reactions or**
 108 **General Adverse Events Within 4 Days of Vaccination * With KINRIX or Separate**
 109 **Concomitant Administration of INFANRIX and IPV When Coadministered With MMR**
 110 **Vaccine (Study 048) (Total Vaccinated Cohort)**

	KINRIX	INFANRIX + IPV
Local[†]	N = 3,121-3,128	N = 1,039-1,043
Pain, any	57.0 [‡]	53.3
Pain, grade 2 or 3 [§]	13.7	12.0
Pain, grade 3 [§]	1.6 [‡]	0.6
Redness, any	36.6	36.6
Redness, ≥50 mm	17.6	20.0
Redness, ≥110 mm	2.9	4.1
Arm circumference increase, any	36.0	37.8
Arm circumference increase, >20 mm	6.9	7.4
Arm circumference increase, >30 mm	2.4	3.2
Swelling, any	26.0	27.0
Swelling, ≥50 mm	10.2	11.5
Swelling, ≥110 mm	1.4	1.8
General	N = 3,037-3,120	N = 993-1,036
Drowsiness, any	19.1	17.5
Drowsiness, grade 3	0.8	0.8
Fever, ≥99.5°F	16.0	14.8
Fever, >100.4°F	6.5 [‡]	4.4
Fever, >102.2°F	1.1	1.1
Fever, >104°F	0.1	0.0
Loss of appetite, any	15.5	16.0
Loss of appetite, grade 3 [¶]	0.8	0.6

111 IPV manufactured by Sanofi Pasteur SA. MMR vaccine manufactured by Merck & Co., Inc.

112 Total Vaccinated Cohort = all vaccinated subjects for whom safety data were available.

113 N = number of children with evaluable data for the events listed.

114 * Within 4 days of vaccination defined as day of vaccination and the next 3 days.

115 † Local reactions at the injection site for KINRIX or INFANRIX.

116 ‡ Statistically higher than comparator group (p<0.05).

117 § Grade 2 defined as painful when the limb was moved; Grade 3 defined as preventing normal
 118 daily activities.

119 || Grade 3 defined as preventing normal daily activities.

120 ¶ Grade 3 defined as not eating at all.

121

122 In Study 048, KINRIX was non-inferior to INFANRIX with regard to swelling that
 123 involved >50% of the injected upper arm length and that was associated with a >30 mm increase
 124 in mid-upper arm circumference within 4 days following vaccination (upper limit of two-sided

125 95% Confidence Interval for difference in percentage of KINRIX [0.6%, n = 20] minus
126 INFANRIX [1.0%, n = 11] $\leq 2\%$).

127 Serious Adverse Events: Within the 31-day period following study vaccination in 3
128 studies (Study 046, 047, and 048), in which all subjects received concomitant MMR vaccine (US
129 licensed MMR vaccine [Merck & Co., Inc.] in Study 047 and 048; non-US licensed MMR
130 vaccine in Study 046), 3 subjects (0.1% [3/3,537]) who received KINRIX reported serious
131 adverse events (dehydration and hypernatremia; cerebrovascular accident; dehydration and
132 gastroenteritis) and 4 subjects (0.3% [4/1,434]) who received INFANRIX and IPV (Sanofi
133 Pasteur SA) reported serious adverse events (cellulitis; constipation; foreign body trauma; fever
134 without identified etiology).

135 **6.2 Postmarketing Experience**

136 In addition to reports in clinical trials, the following adverse events, for which a causal
137 relationship to components of KINRIX is plausible, have been reported since market
138 introduction of DTaP-IPV manufactured by GlaxoSmithKline outside the U.S. Because these
139 events are reported voluntarily from a population of uncertain size, it is not always possible to
140 reliably estimate their frequency or establish a causal relationship to vaccination.

141 General Disorders and Administration Site Conditions: Injection site vesicles.

142 Skin and Subcutaneous Tissue Disorders: Pruritus.

143 Additional adverse events reported following postmarketing use of INFANRIX, for
144 which a causal relationship to vaccination is plausible, are: Allergic reactions, including
145 anaphylactoid reactions, anaphylaxis, angioedema, and urticaria, apnea, collapse or shock-like
146 state (hypotonic-hyporesponsive episode), convulsions (with or without fever),
147 lymphadenopathy, and thrombocytopenia.

148 **7 DRUG INTERACTIONS**

149 **7.1 Concomitant Vaccine Administration**

150 In clinical trials, KINRIX was administered concomitantly with the second dose of MMR
151 vaccine [*see Clinical Studies (14)*].

152 Data are not available on concomitant use of KINRIX and varicella vaccine.

153 When KINRIX is administered concomitantly with other injectable vaccines, they should
154 be given with separate syringes. KINRIX should not be mixed with any other vaccine in the
155 same syringe or vial.

156 **7.2 Immunosuppressive Therapies**

157 Immunosuppressive therapies, including irradiation, antimetabolites, alkylating agents,
158 cytotoxic drugs, and corticosteroids (used in greater than physiologic doses), may reduce the
159 immune response to KINRIX.

160 **8 USE IN SPECIFIC POPULATIONS**

161 **8.1 Pregnancy**

162 Pregnancy Category C

163 Animal reproduction studies have not been conducted with KINRIX. It is also not known

164 whether KINRIX can cause fetal harm when administered to a pregnant woman or can affect
165 reproduction capacity.

166 **8.4 Pediatric Use**

167 Safety and effectiveness of KINRIX in children younger than 4 years of age and children
168 7 to 16 years of age have not been evaluated. KINRIX is not approved for use in persons in these
169 age groups.

170 **11 DESCRIPTION**

171 KINRIX (Diphtheria and Tetanus Toxoids and Acellular Pertussis Adsorbed and
172 Inactivated Poliovirus Vaccine) is a noninfectious, sterile vaccine for intramuscular
173 administration. Each 0.5-mL dose is formulated to contain 25 Lf of diphtheria toxoid, 10 Lf of
174 tetanus toxoid, 25 mcg of inactivated pertussin toxin (PT), 25 mcg of filamentous hemagglutinin
175 (FHA), 8 mcg of pertactin (69 kiloDalton outer membrane protein), 40 D-antigen Units (DU) of
176 Type 1 poliovirus (Mahoney), 8 DU of Type 2 poliovirus (MEF-1), and 32 DU of Type 3
177 poliovirus (Saukett). The diphtheria, tetanus, and pertussis components of KINRIX are the same
178 as those in INFANRIX and PEDIARIX and the poliovirus component is the same as that in
179 PEDIARIX.

180 The diphtheria toxin is produced by growing *Corynebacterium diphtheriae* in Fenton
181 medium containing a bovine extract. Tetanus toxin is produced by growing *Clostridium tetani* in
182 a modified Latham medium derived from bovine casein. The bovine materials used in these
183 extracts are sourced from countries which the United States Department of Agriculture (USDA)
184 has determined neither have nor are at risk of bovine spongiform encephalopathy (BSE). Both
185 toxins are detoxified with formaldehyde, concentrated by ultrafiltration, and purified by
186 precipitation, dialysis, and sterile filtration.

187 The acellular pertussis antigens (PT, FHA, and pertactin) are isolated from *Bordetella*
188 *pertussis* culture grown in modified Stainer-Scholte liquid medium. PT and FHA are isolated
189 from the fermentation broth; pertactin is extracted from the cells by heat treatment and
190 flocculation. The antigens are purified in successive chromatographic and precipitation steps. PT
191 is detoxified using glutaraldehyde and formaldehyde. FHA and pertactin are treated with
192 formaldehyde.

193 Diphtheria and tetanus toxoids and pertussis antigens (inactivated PT, FHA, and
194 pertactin) are individually adsorbed onto aluminum hydroxide.

195 The inactivated poliovirus component of KINRIX is an enhanced potency component.
196 Each of the 3 strains of poliovirus is individually grown in VERO cells, a continuous line of
197 monkey kidney cells, cultivated on microcarriers. Calf serum and lactalbumin hydrolysate are
198 used during VERO cell culture and/or virus culture. Calf serum is sourced from countries the
199 USDA has determined neither have nor are at risk of BSE. After clarification, each viral
200 suspension is purified by ultrafiltration, diafiltration, and successive chromatographic steps, and
201 inactivated with formaldehyde. The 3 purified viral strains are then pooled to form a trivalent
202 concentrate.

203 Diphtheria and tetanus toxoid potency is determined by measuring the amount of
204 neutralizing antitoxin in previously immunized guinea pigs. The potency of the acellular
205 pertussis components (inactivated PT, FHA, and pertactin) is determined by enzyme-linked
206 immunosorbent assay (ELISA) on sera from previously immunized mice. The potency of the
207 inactivated poliovirus component is determined by using the D-antigen ELISA and by a
208 poliovirus neutralizing cell culture assay on sera from previously immunized rats.

209 Each 0.5-mL dose contains 4.5 mg of NaCl and aluminum adjuvant (not more than
210 0.6 mg aluminum by assay). Each dose also contains ≤ 100 mcg of residual formaldehyde and
211 ≤ 100 mcg of polysorbate 80 (Tween 80). Neomycin sulfate and polymyxin B are used in the
212 poliovirus vaccine manufacturing process and may be present in the final vaccine at ≤ 0.05 ng
213 neomycin and ≤ 0.01 ng polymyxin B per dose.

214 KINRIX does not contain a preservative.

215 **12 CLINICAL PHARMACOLOGY**

216 **12.1 Mechanism of Action**

217 Diphtheria: Diphtheria is an acute toxin-mediated infectious disease caused by toxigenic
218 strains of *C. diphtheriae*. Protection against disease is due to the development of neutralizing
219 antibodies to the diphtheria toxin. A serum diphtheria antitoxin level of 0.01 IU/mL is the lowest
220 level giving some degree of protection; a level of 0.1 IU/mL is regarded as protective.¹

221 Tetanus: Tetanus is an acute toxin-mediated disease caused by a potent exotoxin
222 released by *C. tetani*. Protection against disease is due to the development of neutralizing
223 antibodies to the tetanus toxin. A serum tetanus antitoxin level of at least 0.01 IU/mL, measured
224 by neutralization assays, is considered the minimum protective level.^{2,3} A level of ≥ 0.1 IU/mL is
225 considered protective.⁴

226 Pertussis: Pertussis (whooping cough) is a disease of the respiratory tract caused by *B.*
227 *pertussis*. The role of the different components produced by *B. pertussis* in either the
228 pathogenesis of, or the immunity to, pertussis is not well understood. There is no well established
229 serological correlate of protection for pertussis. The efficacy of the pertussis component of
230 KINRIX was determined in clinical trials of INFANRIX administered as a 3-dose series in
231 infants (see INFANRIX prescribing information).

232 Poliomyelitis: Poliovirus is an enterovirus that belongs to the picornavirus family. Three
233 serotypes of poliovirus have been identified (Types 1, 2, and 3). Neutralizing antibodies against
234 the 3 poliovirus serotypes are recognized as conferring protection against poliomyelitis disease.⁵

235 **13 NONCLINICAL TOXICOLOGY**

236 **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

237 KINRIX has not been evaluated for carcinogenic or mutagenic potential, or for
238 impairment of fertility.

239 **14 CLINICAL STUDIES**

240 **14.1 Immunological Evaluation**

241 In a US multicenter study (Study 048), 4,209 children were randomized in a 3:1 ratio to
242 receive either KINRIX or INFANRIX and IPV (Sanofi Pasteur SA) administered concomitantly
243 at separate sites. Subjects also received MMR vaccine (Merck & Co., Inc.) administered
244 concomitantly at a separate site. Subjects were children 4 through 6 years of age who previously
245 received 4 doses of INFANRIX, 3 doses of IPV, and 1 dose of MMR vaccine. Among subjects in
246 both vaccine groups combined, 49.6% were female; 45.6% of subjects were White, 18.8%
247 Hispanic, 13.6% Asian, 7.0% Black, and 15.0% were of other racial/ethnic groups.

248 Levels of antibodies to the diphtheria, tetanus, pertussis (PT, FHA, and pertactin), and
249 poliovirus antigens were measured in sera obtained immediately prior to vaccination and
250 1 month (range 31 to 48 days) after vaccination (Table 2). The co-primary immunogenicity
251 endpoints were anti-diphtheria toxoid, anti-tetanus toxoid, anti-PT, anti-FHA, and anti-pertactin
252 booster responses, and anti-poliovirus Type 1, Type 2, and Type 3 geometric mean antibody
253 titers (GMTs) 1 month after vaccination. KINRIX was shown to be non-inferior to INFANRIX
254 and IPV administered separately, in terms of booster responses to DTaP antigens and post-
255 vaccination GMTs for anti-poliovirus antibodies (Table 2).

256

257 **Table 2. Pre-vaccination Antibody Levels and Post-vaccination* Antibody Responses**
 258 **Following KINRIX Compared With Separate Concomitant Administration of INFANRIX**
 259 **and IPV in Children 4 to 6 Years of Age When Coadministered With MMR Vaccine (Study**
 260 **048) (ATP Cohort for Immunogenicity)**

	KINRIX	INFANRIX + IPV
	N = 787-851	N = 237-262
Anti-Diphtheria Toxoid		
Pre-vaccination % ≥ 0.1 IU/mL (95% CI) [†]	87.7 (85.3, 89.9)	85.5 (80.6, 89.5)
Post-vaccination % ≥ 0.1 IU/mL (95% CI) [†]	100 (99.6, 100)	100 (98.6, 100)
% Booster Response (95% CI) [‡]	99.5 (98.8, 99.9) [§]	100 (98.6, 100)
Anti-Tetanus Toxoid		
Pre-vaccination % ≥ 0.1 IU/mL (95% CI) [†]	87.8 (85.4, 90.0)	88.2 (83.6, 91.8)
Post-vaccination % ≥ 0.1 IU/mL (95% CI) [†]	100 (99.6, 100)	100 (98.6, 100)
% Booster Response (95% CI) [‡]	96.7 (95.2, 97.8) [§]	93.9 (90.2, 96.5)
Anti-PT		
% Booster Response (95% CI)	92.2 (90.2, 94.0) [§]	92.6 (88.7, 95.5)
Anti-FHA		
% Booster Response (95% CI)	95.4 (93.7, 96.7) [§]	96.2 (93.1, 98.1)
Anti-Pertactin		
% Booster Response (95% CI)	97.8 (96.5, 98.6) [§]	96.9 (94.1, 98.7)
Anti-Poliovirus 1		
Pre-vaccination % $\geq 1:8$ (95% CI) [†]	88.3 (85.9, 90.4)	85.1 (80.1, 89.2)
Post-vaccination % $\geq 1:8$ (95% CI) [†]	99.9 (99.3, 100)	100 (98.5, 100)
Post-vaccination GMT (95% CI)	2,127 (1,976, 2,290) [¶]	1,685 (1,475, 1,925)
Anti-Poliovirus 2		
Pre-vaccination % $\geq 1:8$ (95% CI) [†]	91.8 (89.7, 93.6)	87.0 (82.3, 90.8)
Post-vaccination % $\geq 1:8$ (95% CI) [†]	100 (99.6, 100)	100 (98.5, 100)
Post-vaccination GMT (95% CI)	2,265 (2,114, 2,427) [¶]	1,818 (1,606, 2,057)
Anti-Poliovirus 3		
Pre-vaccination % $\geq 1:8$ (95% CI) [†]	84.7 (82.0, 87.0)	85.0 (80.1, 89.1)
Post-vaccination % $\geq 1:8$ (95% CI) [†]	100 (99.5, 100)	100 (98.5, 100)
Post-vaccination GMT (95% CI)	3,588 (3,345, 3,849) [¶]	3,365 (2,961, 3,824)

261 IPV manufactured by Sanofi Pasteur SA. MMR vaccine manufactured by Merck & Co., Inc.

262 ATP = according-to-protocol; CI = Confidence Interval; GMT = geometric mean antibody titer

263 N = number of subjects with available results.

264 * One month blood sampling, range 31 to 48 days.

265 † Seroprotection defined as anti-diphtheria toxoid and anti-tetanus toxoid antibody
 266 concentrations ≥ 0.1 IU/mL by ELISA and as anti-poliovirus Type 1, Type 2, and Type 3
 267 antibody titer $\geq 1:8$ by micro-neutralization assay for poliovirus.

268 ‡ Booster response: In subjects with pre-vaccination < 0.1 IU/mL, post-vaccination
 269 concentration ≥ 0.4 IU/mL. In subjects with pre-vaccination concentration ≥ 0.1 IU/mL, an
 270 increase of at least 4 times the pre-vaccination concentration.

271 § KINRIX was non-inferior to INFANRIX + IPV based on booster response rates (upper limit

272 of two-sided 95% CI on the difference of INFANRIX + IPV minus KINRIX $\leq 10\%$).
273 ¶ Booster response: In subjects with pre-vaccination < 5 EL.U./mL, post-vaccination
274 concentration ≥ 20 EL.U./mL. In subjects with pre-vaccination ≥ 5 EL.U./mL and
275 < 20 EL.U./mL, an increase of at least 4 times the pre-vaccination concentration. In subjects
276 with pre-vaccination ≥ 20 EL.U./mL, an increase of at least 2 times the pre-vaccination
277 concentration.
278 ¶ KINRIX was non-inferior to INFANRIX + IPV based on post-vaccination anti-poliovirus
279 antibody GMTs adjusted for baseline titer (upper limit of two-sided 95% CI for the GMT ratio
280 [INFANRIX + IPV:KINRIX] ≤ 1.5).

281
282 **14.2 Concomitant Vaccine Administration**
283 In a US study (Study 047), among recipients of DTaP-IPV (same formulation as KINRIX
284 but also containing 2-phenoxyethanol) and the second dose of MMR vaccine (Merck & Co.,
285 Inc.) who had pre-vaccination sera tested for antibodies to measles, mumps, and rubella
286 (N = 175-181), 99% of subjects were seropositive for antibodies to measles, mumps, and rubella
287 prior to vaccination.

288 **15 REFERENCES**

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304 **16 HOW SUPPLIED/STORAGE AND HANDLING**

305 KINRIX is available in 0.5-mL single-dose vials and disposable prefilled TIP-LOK
306 syringes.

307 Single-Dose Vials

308 NDC 58160-812-11 (package of 10)

309 Single-Dose Prefilled Disposable TIP-LOK Syringes (packaged without needles)

310 NDC 58160-812-46 (package of 5)

311 Store refrigerated between 2° and 8°C (36° and 46°F). Do not freeze. Discard if the
312 vaccine has been frozen.

313 **17 PATIENT COUNSELING INFORMATION**

- 314 • Parents or guardians should be informed by the healthcare provider of the potential benefits
315 and risks of immunization with KINRIX.
- 316 • The healthcare provider should inform the parents or guardians about the potential for
317 adverse reactions that have been temporally associated with administration of KINRIX or
318 other vaccines containing similar components.
- 319 • The parent or guardian accompanying the recipient should be instructed to report any adverse
320 events to their healthcare provider where the vaccine was administered.
- 321 • The parent or guardian should be given the Vaccine Information Statements, which are
322 required by the National Childhood Vaccine Injury Act of 1986 to be given prior to
323 immunization. These materials are available free of charge at the Centers for Disease Control
324 and Prevention (CDC) website (www.cdc.gov/nip).

325
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327 GlaxoSmithKline. IPOL is a registered trademark of Sanofi Pasteur Limited.

328



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