HIGHLIGHTS OF PRESCRIBING INFORMATION

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

BOOSTRIX (Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine, Adsorbed) Suspension for Intramuscular Injection

Initial U.S. Approval: 2005

RECENT MAJOR CHANGES		
Indications and Usage (1)	07/2011	
Warnings and Precautions, Syncope (5.3)	03/2012	
INDICATIONS AND USAGE		

BOOSTRIX is a vaccine indicated for active booster immunization against tetanus, diphtheria, and pertussis. BOOSTRIX is approved for use as a single dose in individuals 10 years of age and older. (1)

----- DOSAGE AND ADMINISTRATION ------A single intramuscular injection (0.5 mL). (2.2)

----- DOSAGE FORMS AND STRENGTHS ------Single-dose vials and prefilled syringes containing a 0.5-mL suspension for injection. (3)

--CONTRAINDICATIONS--

- Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any tetanus toxoid-, diphtheria toxoid-, or pertussis antigen-containing vaccine or to any component of BOOSTRIX. (4.1)
- Encephalopathy (e.g., coma, decreased level of consciousness, prolonged seizures) within 7 days of administration of a previous pertussis antigen-containing vaccine. (4.2)

- WARNINGS AND PRECAUTIONS--

- BOOSTRIX is available in vials and 2 types of prefilled syringes. One type of prefilled syringe has a tip cap which may contain natural rubber latex. The other type has a tip cap and a rubber plunger which contain dry natural latex rubber. Use of these syringes may cause allergic reactions in latex-sensitive individuals. (5.1, 16)
- If Guillain-Barré syndrome occurred within 6 weeks of receipt of a prior vaccine containing tetanus toxoid, the risk of Guillain-Barré syndrome may be increased following a subsequent dose of tetanus toxoidcontaining vaccine, including BOOSTRIX. (5.2)
- Syncope (fainting) can occur in association with administration of injectable vaccines, including BOOSTRIX. Procedures should be in place to avoid falling injury and to restore cerebral perfusion following syncope. (5.3)

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- Progressive or unstable neurologic conditions are reasons to defer vaccination with a pertussis-containing vaccine, including BOOSTRIX. (5.4)
- Persons who experienced an Arthus-type hypersensitivity reaction following a prior dose of a tetanus toxoid-containing vaccine should not receive BOOSTRIX unless at least 10 years have elapsed since the last dose of a tetanus toxoid-containing vaccine. (5.5)

----- ADVERSE REACTIONS ------

- Common solicited adverse events (≥15%) in adolescents (10 to 18 years • of age) were pain, redness, and swelling at the injection site, increase in arm circumference of injected arm, headache, fatigue, and gastrointestinal symptoms. (6.1)
- Common solicited adverse events (≥15%) in adults (19 to 64 years of age) were pain, redness, and swelling at the injection site, headache, fatigue, and gastrointestinal symptoms. (6.1)
- The most common solicited adverse event ($\geq 15\%$) in the elderly (65 years of age and older) was pain at the injection site. (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 ------
- In subjects 11 to 18 years of age, lower levels for antibodies to pertactin were observed when BOOSTRIX was administered concomitantly with meningococcal conjugate vaccine (serogroups A, C, Y, and W-135) as compared to BOOSTRIX administered first. (7.1)
- In subjects 19 to 64 years of age, lower levels for antibodies to FHA and pertactin were observed when BOOSTRIX was administered concomitantly with an inactivated influenza vaccine as compared to BOOSTRIX alone. (7.1)
- Do not mix BOOSTRIX with any other vaccine in the same syringe or vial. (7.1)

------ USE IN SPECIFIC POPULATIONS ------

- Safety and effectiveness of BOOSTRIX have not been established in pregnant women. (8.1)
- Register women who receive BOOSTRIX while pregnant in the pregnancy registry by calling 1-888-452-9622. (8.1)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: XX/XXXX

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1 FULL PRESCRIBING INFORMATION

2 1 INDICATIONS AND USAGE

BOOSTRIX[®] is indicated for active booster immunization against tetanus, diphtheria,
and pertussis. BOOSTRIX is approved for use as a single dose in individuals 10 years of age and
older.

6 2 DOSAGE AND ADMINISTRATION

7 2.1 Preparation for Administration

8 Shake vigorously to obtain a homogeneous, turbid, white suspension before

9 administration. Do not use if resuspension does not occur with vigorous shaking. Parenteral drug

10 products should be inspected visually for particulate matter and discoloration prior to

11 administration, whenever solution and container permit. If either of these conditions exists, the

12 vaccine should not be administered.

For the prefilled syringes, attach a sterile needle and administer intramuscularly.

14 For the vials, use a sterile needle and sterile syringe to withdraw the 0.5-mL dose and

15 administer intramuscularly. Changing needles between drawing vaccine from a vial and injecting

16 it into a recipient is not necessary unless the needle has been damaged or contaminated. Use a

- 17 separate sterile needle and syringe for each individual.
- 18 Do not administer this product intravenously, intradermally, or subcutaneously.

19 **2.2 Dose and Schedule**

- 20 BOOSTRIX is administered as a single 0.5-mL intramuscular injection into the deltoid 21 muscle of the upper arm.
- 22 There are no data to support repeat administration of BOOSTRIX.
- 23 Five years should elapse between the last dose of the recommended series of Diphtheria

24 and Tetanus Toxoids and Acellular Pertussis Vaccine Adsorbed (DTaP) and/or Tetanus and

25 Diphtheria Toxoids Adsorbed For Adult Use (Td) vaccine and the administration of

26 BOOSTRIX.

13

27 2.3 Additional Dosing Information

- 28 <u>Primary Series:</u> The use of BOOSTRIX as a primary series or to complete the primary
 29 series for diphtheria, tetanus, or pertussis has not been studied.
- 30 <u>Wound Management:</u> If tetanus prophylaxis is needed for wound management,
- 31 BOOSTRIX may be given if no previous dose of any Tetanus Toxoid, Reduced Diphtheria

32 Toxoid and Acellular Pertussis Vaccine, Adsorbed (Tdap) has been administered.

33 3 DOSAGE FORMS AND STRENGTHS

- 34 BOOSTRIX is a suspension for injection available in 0.5-mL single-dose vials and
- 35 prefilled TIP-LOK[®] syringes.

36 4 CONTRAINDICATIONS

37 4.1 Hypersensitivity

A severe allergic reaction (e.g., anaphylaxis) after a previous dose of any tetanus toxoid-, diphtheria toxoid-, or pertussis antigen-containing vaccine or any component of this vaccine is a contraindication to administration of BOOSTRIX *[see Description (11)]*. Because of the uncertainty as to which component of the vaccine might be responsible, none of the components should be administered. Alternatively, such individuals may be referred to an allergist for evaluation if immunization with any of these components is considered.

44 4.2 Encephalopathy

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

49 5 WARNINGS AND PRECAUTIONS

50 **5.1 Latex**

51 BOOSTRIX is available in vials and 2 types of prefilled syringes. One type of prefilled 52 syringe has a tip cap which may contain natural rubber latex and a plunger which does not 53 contain latex. The other type has a tip cap and a rubber plunger which contain dry natural latex 54 rubber. Use of these syringes may cause allergic reactions in latex-sensitive individuals. The vial 55 stopper does not contain latex. *[See How Supplied/Storage and Handling (16).]*

56 5.2 Guillain-Barré Syndrome and Brachial Neuritis

57 If Guillain-Barré syndrome occurred within 6 weeks of receipt of a prior vaccine 58 containing tetanus toxoid, the risk of Guillain-Barré syndrome may be increased following a 59 subsequent dose of tetanus toxoid-containing vaccine, including BOOSTRIX. A review by the 60 Institute of Medicine (IOM) found evidence for a causal relationship between receipt of tetanus 61 toxoid and both brachial neuritis and Guillain-Barré syndrome.¹

62 **5.3 Syncope**

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

67 5.4 Progressive or Unstable Neurologic Disorders

68 Progressive or unstable neurologic conditions (e.g., cerebrovascular events and acute 69 encephalopathic conditions) are reasons to defer vaccination with a pertussis-containing vaccine, 70 including BOOSTRIX. It is not known whether administration of BOOSTRIX to persons with an 71 unstable or progressive neurologic disorder might hasten manifestations of the disorder or affect 72 the prognosis. Administration of BOOSTRIX to persons with an unstable or progressive

73 neurologic disorder may result in diagnostic confusion between manifestations of the underlying

74 illness and possible adverse effects of vaccination.

75 5.5 Arthus-Type Hypersensitivity

Persons who experienced an Arthus-type hypersensitivity reaction following a prior dose
of a tetanus toxoid-containing vaccine usually have a high serum tetanus antitoxin level and
should not receive BOOSTRIX or other tetanus toxoid-containing vaccines unless at least
10 years have elapsed since the last dose of tetanus toxoid-containing vaccine.

- 80 **5.6 Altered Immunocompetence**
- 81 As with any vaccine, if administered to immunosuppressed persons, including individuals 82 receiving immunosuppressive therapy, the expected immune response may not be obtained.

83 5.7 Prevention and Management of Acute Allergic Reactions

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

89 6 ADVERSE REACTIONS

90 6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a vaccine cannot be directly compared to rates in the clinical trials of another vaccine, and may not reflect the rates observed in practice. As with any vaccine, there is the possibility that broad use of BOOSTRIX could reveal adverse reactions not observed in clinical trials.

In clinical studies, 4,949 adolescents (10 to 18 years of age) and 4,076 adults (19 years of
 age and older) were vaccinated with a single dose of BOOSTRIX. Of these adolescents, 1,341

98 were vaccinated with BOOSTRIX in a coadministration study with meningococcal conjugate

vaccine [see Drug Interactions (7.1) and Clinical Studies (14.5)]. Of these adults, 1,104 were

100 65 years of age and older [see Clinical Studies (14.4)]. A total of 860 adults19 years of age and

101 older received concomitant vaccination with BOOSTRIX and influenza vaccines in a

102 coadministration study [see Drug Interactions (7.1) and Clinical Studies (14.5)]. An additional

103 1,092 adolescents 10 to 18 years of age received a non-US formulation of BOOSTRIX

104 (formulated to contain 0.5 mg aluminum per dose) in non-US clinical studies.

105 In a randomized, observer-blinded, controlled study in the US, 3,080 adolescents 10 to 106 18 years of age received a single dose of BOOSTRIX and 1,034 received the comparator Td

107 vaccine, manufactured by MassBioLogics. There were no substantive differences in

108 demographic characteristics between the vaccine groups. Among BOOSTRIX and comparator

vaccine recipients, approximately 75% were 10 to 14 years of age and approximately 25% were

- 110 15 to 18 years of age. Approximately 98% of participants in this study had received the
- recommended series of 4 or 5 doses of either Diphtheria and Tetanus Toxoids and Pertussis
- 112 Vaccine Adsorbed (DTwP) or a combination of DTwP and DTaP in childhood. Subjects were
- 113 monitored for solicited adverse events using standardized diary cards (day 0-14). Unsolicited

adverse events were monitored for the 31-day period following vaccination (day 0-30). Subjects

- 115 were also monitored for 6 months post-vaccination for non-routine medical visits, visits to an
- 116 emergency room, onset of new chronic illness, and serious adverse events. Information regarding
- 117 late onset adverse events was obtained via a telephone call 6 months following vaccination. At
- 118 least 97% of subjects completed the 6-month follow-up evaluation.

119 In a study conducted in Germany, BOOSTRIX was administered to 319 children 10 to 120 12 years of age previously vaccinated with 5 doses of acellular pertussis antigen-containing vaccines: 193 of these subjects had previously received 5 doses of INFANRIX[®] (Diphtheria and 121 122 Tetanus Toxoids and Acellular Pertussis Vaccine Adsorbed). Adverse events were recorded on 123 diary cards during the 15 days following vaccination. Unsolicited adverse events that occurred 124 within 31 days of vaccination (day 0-30) were recorded on the diary card or verbally reported to 125 the investigator. Subjects were monitored for 6 months post-vaccination for physician office 126 visits, emergency room visits, onset of new chronic illness, and serious adverse events. The 6-127 month follow-up evaluation, conducted via telephone interview, was completed by 90% of 128 subjects.

129 The US adult (19 to 64 years of age) study, a randomized, observer-blinded study, 130 evaluated the safety of BOOSTRIX (N = 1,522) compared with ADACEL[®] (Tetanus Toxoid, 121 Delay 1 D

Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine Adsorbed) (N = 762), a Tdap

132 vaccine manufactured by Sanofi Pasteur SA. Vaccines were administered as a single dose. There

133 were no substantive differences in demographic characteristics between the vaccine groups.

134 Subjects were monitored for solicited adverse events using standardized diary cards (day 0-14).

135 Unsolicited adverse events were monitored for the 31-day period following vaccination (day 0-

30). Subjects were also monitored for 6 months post-vaccination for serious adverse events,visits to an emergency room, hospitalizations, and onset of new chronic illness. Approximately

138 95% of subjects completed the 6-month follow-up evaluation.

139 The US elderly (65 years of age and older) study, a randomized, observer-blinded study, evaluated the safety of BOOSTRIX (N = 887) compared with DECAVAC[®] (Tetanus and 140 141 Diphtheria Toxoids Adsorbed) (N = 445), a US-licensed Td vaccine, manufactured by Sanofi 142 Pasteur SA. Vaccines were administered as a single dose. Among all vaccine recipients, the 143 mean age was approximately 72 years; 54% were female and 95% were white. Subjects were 144 monitored for solicited adverse events using standardized diary cards (day 0-3). Unsolicited 145 adverse events were monitored for the 31-day period following vaccination (day 0-30). Subjects 146 were also monitored for 6 months post-vaccination for serious adverse events. Approximately

147 99% of subjects completed the 6-month follow-up evaluation.

- Solicited Adverse Events in the US Adolescent Study: Table 1 presents the solicited
 local adverse reactions and general adverse events within 15 days of vaccination with
 BOOSTRIX or Td vaccine for the total vaccinated cohort
- 150 BOOSTRIX or Td vaccine for the total vaccinated cohort.
- The primary safety endpoint was the incidence of grade 3 pain (spontaneously painful and/or prevented normal activity) at the injection site within 15 days of vaccination. Grade 3 pain was reported in 4.6% of those who received BOOSTRIX compared with 4.0% of those who

- 154 received the Td vaccine. The difference in rate of grade 3 pain was within the pre-defined
- 155 clinical limit for non-inferiority (upper limit of the 95% CI for the difference [BOOSTRIX minus
- 156 Td] ≤4%).
- 157

158	Table 1. Rates of Solicited Loc	al Adverse Reactions of	r General Adverse Events	Within the 15-
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	BOOSTRIX	Td
	(N = 3,032)	(N = 1,013)
	%	%
Local		
Pain, any ^b	75.3	71.7
Pain, grade 2 or 3 ^b	51.2	42.5
Pain, grade 3 ^c	4.6	4.0
Redness, any	22.5	19.8
Redness, >20 mm	4.1	3.9
Redness, ≥50 mm	1.7	1.6
Swelling, any	21.1	20.1
Swelling, >20 mm	5.3	4.9
Swelling, ≥50 mm	2.5	3.2
Arm circumference increase, >5 mm ^d	28.3	29.5
Arm circumference increase, >20 mm ^d	2.0	2.2
Arm circumference increase, >40 mm ^d	0.5	0.3
General		
Headache, any	43.1	41.5
Headache, grade 2 or 3 ^b	15.7	12.7
Headache, grade 3	3.7	2.7
Fatigue, any	37.0	36.7
Fatigue, grade 2 or 3	14.4	12.9
Fatigue, grade 3	3.7	3.2
Gastrointestinal symptoms, any ^e	26.0	25.8
Gastrointestinal symptoms, grade 2 or 3 ^e	9.8	9.7
Gastrointestinal symptoms, grade 3 ^e	3.0	3.2
Fever, $\ge 99.5^{\circ}F (37.5^{\circ}C)^{f}$	13.5	13.1
Fever, $>100.4^{\circ}F(38.0^{\circ}C)^{f}$	5.0	4.7
Fever. $>102.2^{\circ}F(39.0^{\circ}C)^{f}$	1.4	1.0

159 day^a Post-Vaccination Period in Adolescents 10 to 18 Years of Age (Total Vaccinated Cohort)

160 Td = Tetanus and Diphtheria Toxoids Adsorbed For Adult Use manufactured by MassBioLogics.

- 161 N = Number of subjects in the total vaccinated cohort with local/general symptoms sheets
 162 completed.
- 163 Grade 2 = Local: painful when limb moved; General: interfered with normal activity.
- 164 Grade 3 = Local: spontaneously painful and/or prevented normal activity; General: prevented
- 165 normal activity.
- 166 ^a Day of vaccination and the next 14 days.
- ^b Statistically significantly higher (P < 0.05) following BOOSTRIX as compared to Td vaccine.
- 168 ^c Grade 3 injection site pain following BOOSTRIX was not inferior to Td vaccine (upper limit 169 of two-sided 95% CI for the difference [BOOSTRIX minus Td] in the percentage of subjects 170 $\leq 4\%$).
- ^d Mid-upper region of the vaccinated arm.
- 172 ^e Gastrointestinal symptoms included nausea, vomiting, diarrhea, and/or abdominal pain.
- ^f Oral temperatures or axillary temperatures.

- 175 <u>Unsolicited Adverse Events in the US Adolescent Study:</u> The incidence of 176 unsolicited adverse events reported in the 31 days after vaccination was comparable between the
- 2 groups (25.4% and 24.5% for BOOSTRIX and Td vaccine, respectively).
- 178 Solicited Adverse Events in the German Adolescent Study: Table 2 presents the

179 rates of solicited local adverse reactions and fever within 15 days of vaccination for those180 subjects who had previously been vaccinated with 5 doses of INFANRIX. No cases of whole

180 subjects who had previously been vaccinated with 5 doses of INFANRIX. No cases of whole 181 arm swelling were reported. Two individuals (2/193) reported large injection site swelling (range

182 110 to 200 mm diameter), in one case associated with grade 3 pain. Neither individual sought

183 medical attention. These episodes were reported to resolve without sequelae within 5 days.

184

174

185Table 2. Rates of Solicited Adverse Events Reported Within the 15-day^a Post-Vaccination

186 Period Following Administration of BOOSTRIX in Adolescents 10 to 12 Years of Age Who

187 Had Previously Received 5 Doses of INFANRIX

	BOOSTRIX
	(N = 193)
	%
Pain, any	62.2
Pain, grade 2 or 3	33.2
Pain, grade 3	5.7
Redness, any	47.7
Redness, >20 mm	15.0
Redness, ≥50 mm	10.9
Swelling, any	38.9
Swelling, >20 mm	17.6
Swelling, ≥50 mm	14.0
Fever, ≥99.5°F (37.5°C) ^b	8.8
Fever, >100.4°F (38.0°C) ^b	4.1
Fever, >102.2°F (39.0°C) ^b	1.0

- 188 N = Number of subjects with local/general symptoms sheets completed.
- 189 Grade 2 = Painful when limb moved.
- 190 Grade 3 = Spontaneously painful and/or prevented normal activity.
- ^a Day of vaccination and the next 14 days.
- ^b Oral temperatures or axillary temperatures.
- 193
- 194 Solicited Adverse Events in the US Adult (19 to 64 Years of Age) Study: Table 3
- 195 presents solicited local adverse reactions and general adverse events within 15 days of
- 196 vaccination with BOOSTRIX or the comparator Tdap vaccine for the total vaccinated cohort.
- 197

198	Table 3. Rates of Solicited Local Adverse Reactions or General Adverse Events Within the

	BOOSTRIX Tda	
	(N = 1,480)	(N = 741)
	%	%
Local		
Pain, any	61.0	69.2
Pain, grade 2 or 3	35.1	44.4
Pain, grade 3	1.6	2.3
Redness, any	21.1	27.1
Redness, >20 mm	4.0	6.2
Redness, ≥50 mm	1.6	2.3
Swelling, any	17.6	25.6
Swelling, >20 mm	3.9	6.3
Swelling, ≥50 mm	1.4	2.8
General		
Headache, any	30.1	31.0
Headache, grade 2 or 3	11.1	10.5
Headache, grade 3	2.2	1.5
Fatigue, any	28.1	28.9
Fatigue, grade 2 or 3	9.1	9.4
Fatigue, grade 3	2.5	1.2
Gastrointestinal symptoms, any ^b	15.9	17.5
Gastrointestinal symptoms, grade 2 or 3 ^b	4.3	5.7
Gastrointestinal symptoms, grade 3 ^b	1.2	1.3
Fever, $\ge 99.5^{\circ}F(37.5^{\circ}C)^{\circ}$	5.5	8.0
Fever, $>100.4^{\circ}F(38.0^{\circ}C)^{\circ}$	1.0	1.5
Fever, $>102.2^{\circ}F(39.0^{\circ}C)^{\circ}$	0.1	0.4

199 **15-day^a** Post-Vaccination Period in Adults 19 to 64 Years of Age (Total Vaccinated Cohort)

- 204 Grade 2 = Local: painful when limb moved; General: interfered with normal activity.
- 205 Grade 3 = Local/General: prevented normal activity.
- ^a Day of vaccination and the next 14 days.
- ^b Gastrointestinal symptoms included nausea, vomiting, diarrhea, and/or abdominal pain.
- 208 ^c Oral temperatures.
- 209
- 210 Unsolicited Adverse Events in the US Adult (19 to 64 Years of Age) Study: The
- 211 incidence of unsolicited adverse events reported in the 31 days after vaccination was comparable
- between the 2 groups (17.8% and 22.2% for BOOSTRIX and Tdap vaccine, respectively).
- 213 Solicited Adverse Events in the US Elderly (65 Years of Age and Older) Study:
- Table 4 presents solicited local adverse reactions and general adverse events within 4 days of

Tdap = Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine Adsorbed,
 a Tdap vaccine manufactured by Sanofi Pasteur SA.

N = Number of subjects in the total vaccinated cohort with local/general symptoms sheets
 completed.

- 215 vaccination with BOOSTRIX or the comparator Td vaccine for the total vaccinated cohort.
- 216

217 Table 4. Rates of Solicited Local Adverse Reactions or General Adverse Events Within

218 <u>4 Days^a of Vaccination in the Elderly 65 Years of Age and Older (Total Vaccinated Cohort)</u>

	BOOSTRIX	Td
	%	%
Local	(N = 882)	(N = 444)
Pain, any	21.5	27.7
Pain, grade 2 or 3	7.5	10.1
Pain, grade 3	0.2	0.7
Redness, any	10.8	12.6
Redness, >20 mm	1.4	2.5
Redness, ≥50 mm	0.6	0.9
Swelling, any	7.5	11.7
Swelling, >20 mm	2.2	3.4
Swelling, ≥50 mm	0.7	0.7
General	(N = 882)	(N = 445)
Fatigue, any	12.5	14.8
Fatigue, grade 2 or 3	2.5	2.9
Fatigue, grade 3	0.7	0.7
Headache, any	11.5	11.7
Headache, grade 2 or 3	1.9	2.2
Headache, grade 3	0.6	0.0
Gastrointestinal symptoms, any ^b	7.6	9.2
Gastrointestinal symptoms, grade 2 or 3 ^b	1.7	1.8
Gastrointestinal symptoms, grade 3 ^b	0.3	0.4
Fever, ≥99.5°F $(37.5°C)^{c}$	2.0	2.5
Fever, $>100.4^{\circ}F (38.0^{\circ}C)^{\circ}$	0.2	0.2
Fever, $>102.2^{\circ}F(39.0^{\circ}C)^{\circ}$	0.0	0.0

- Td = Tetanus and Diphtheria Toxoids Adsorbed, a US-licensed Td vaccine, manufactured by
 Sanofi Pasteur SA.
- 221 N = Number of subjects with a documented dose.
- 222 Grade 2 = Local: painful when limb moved; General: interfered with normal activity.
- 223 Grade 3 = Local/General: prevented normal activity.
- ^a Day of vaccination and the next 3 days.
- ^b Gastrointestinal symptoms included nausea, vomiting, diarrhea, and/or abdominal pain.
- ^c Oral temperatures.
- 227
- 228 Unsolicited Adverse Events in the US Elderly (65 Years of Age and Older) Study:

229 The incidence of unsolicited adverse events reported in the 31 days after vaccination was

- comparable between the 2 groups (17.1% and 14.4% for BOOSTRIX and Td vaccine,
- 231 respectively).
- 232 Serious Adverse Events (SAEs): In the US and German adolescent safety studies, no

233 serious adverse events were reported to occur within 31 days of vaccination. During the 6-month 234 extended safety evaluation period, no serious adverse events that were of potential autoimmune 235 origin or new onset and chronic in nature were reported to occur. In non-US adolescent studies in 236 which serious adverse events were monitored for up to 37 days, one subject was diagnosed with 237 insulin-dependent diabetes 20 days following administration of BOOSTRIX. No other serious 238 adverse events of potential autoimmune origin or that were new onset and chronic in nature were 239 reported to occur in these studies. In the US adult (19 to 64 years of age) study, serious adverse 240 events were reported to occur during the entire study period (0-6 months) by 1.4% and 1.7% of 241 subjects who received BOOSTRIX and the comparator Tdap vaccine, respectively. During the 6-242 month extended safety evaluation period, no serious adverse events of a neuroinflammatory 243 nature or with information suggesting an autoimmune etiology were reported in subjects who 244 received BOOSTRIX. In the US elderly (65 years of age and older) study, serious adverse events 245 were reported to occur by 0.7% and 0.9% of subjects who received BOOSTRIX and the 246 comparator Td vaccine, respectively, during the 31-day period after vaccination. Serious adverse events were reported to occur by 4.2% and 2.2% of subjects who received BOOSTRIX and the 247 248 comparator Td vaccine, respectively, during the 6-month period after vaccination. 249 Concomitant Vaccination With Meningococcal Conjugate Vaccine in 250 Adolescents: In a randomized study in the US, 1,341 adolescents (11 to 18 years of age) received either BOOSTRIX administered concomitantly with MENACTRA® (Meningococcal 251 252 (Groups A, C, Y, and W-135) Polysaccharide Diphtheria Toxoid Conjugate Vaccine), (Sanofi 253 Pasteur SA), or each vaccine administered separately 1 month apart [see Drug Interactions (7.1)] 254 and Clinical Studies (14.5)]. Safety was evaluated in 446 subjects who received BOOSTRIX 255 administered concomitantly with meningococcal conjugate vaccine at different injection sites, 256 446 subjects who received BOOSTRIX followed by meningococcal conjugate vaccine 1 month 257 later, and 449 subjects who received meningococcal conjugate vaccine followed by BOOSTRIX 258 1 month later. Solicited local adverse reactions and general adverse events were recorded on 259 diary cards for 4 days (day 0-3) following each vaccination. Unsolicited adverse events were 260 monitored for the 31-day period following each vaccination (day 0-30). Table 5 presents the 261 percentages of subjects experiencing local reactions at the injection site for BOOSTRIX and 262 solicited general events following BOOSTRIX. The incidence of unsolicited adverse events 263 reported in the 31 days after any vaccination was similar following each dose of BOOSTRIX in 264 all cohorts.

266 **Table 5. Rates of Solicited Local Adverse Reactions or General Adverse Events Reported**

267	Within the 4-day Post-V	accination Period following	Administration of BOOSTRIX in
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	$BOOSTRIX+MCV4^{a}$ (N = 441) %	BOOSTRIX \rightarrow MCV4 ^b (N = 432-433) %	$MCV4 \rightarrow BOOSTRIX^{c}$ $(N = 441)$ %
Local (at injection	n site for BOOSTRIX)		
Pain, any	70.1	70.4	47.8
Redness, any	22.7	25.7	17.9
Swelling, any	17.7	18.1	12.0
General (followin	g administration of BOO	STRIX)	
Fatigue	34.0	32.1	20.4
Headache	34.0	30.7	17.0
Gastrointestinal symptoms ^d	15.2	14.5	7.7
Fever, ≥99.5°F (37.5°C) ^e	5.2	3.5	2.3

268 Individuals 11 to 18 Years of Age (Total Vaccinated Cohort)

269	MCV4 = MENACTRA (Meningococcal (Groups A, C, Y, and W-135) Polysaccharide

270 Diphtheria Toxoid Conjugate Vaccine), Sanofi Pasteur SA.

N = number of subjects in the total vaccinated cohort with local/general symptoms sheets
 completed.

^a BOOSTRIX+MCV4 = concomitant vaccination with BOOSTRIX and MENACTRA.

^b BOOSTRIX \rightarrow MCV4 = BOOSTRIX followed by MCV4 1 month later.

275 ^c MCV4 \rightarrow BOOSTRIX = MCV4 followed by BOOSTRIX 1 month later.

^d Gastrointestinal symptoms included nausea, vomiting, diarrhea, and/or abdominal pain.

277 ^e Oral temperatures.

278

279 6.2 Postmarketing Experience

In addition to reports in clinical trials, worldwide voluntary reports of adverse events received for BOOSTRIX in persons 10 years of age and older since market introduction of this vaccine are listed below. This list includes serious events or events which have causal connection to components of this or other vaccines or drugs. Because these events are reported voluntarily from a population of uncertain size, it is not possible to reliably estimate their frequency or establish a causal relationship to the vaccine.

- 286 Blood and Lymphatic System Disorders: Lymphadenitis, lymphadenopathy.
- 287 <u>Cardiac Disorders:</u> Myocarditis.

288 General Disorders and Administration Site Conditions: Extensive swelling of the

289 injected limb, injection site induration, injection site inflammation, injection site mass, injection

- site pruritus, injection site nodule, injection site warmth, local reaction.
- 291 <u>Musculoskeletal and Connective Tissue Disorders:</u> Arthralgia, back pain, myalgia.

- 292 Nervous System Disorders: Convulsion, encephalitis, facial palsy, paraesthesia,
- syncope.

294 <u>Skin and Subcutaneous Tissue Disorders:</u> Exanthem, Henoch-Schönlein purpura,
 295 rash, urticaria.

2967DRUG INTERACTIONS

297 7.1 Concomitant Vaccine Administration

BOOSTRIX was administered concomitantly with MENACTRA in a clinical study of subjects 11 to 18 years of age *[see Clinical Studies (14.5)]*. Post-vaccination geometric mean antibody concentrations (GMCs) to pertactin were lower following BOOSTRIX administered concomitantly with meningococcal conjugate vaccine compared to BOOSTRIX administered first. It is not known if the efficacy of BOOSTRIX is affected by the reduced response to pertactin.

BOOSTRIX was administered concomitantly with FLUARIX[®] (Influenza Virus Vaccine) in a clinical study of subjects 19 to 64 years of age *[see Clinical Studies (14.5)]*. Lower GMCs for antibodies to the pertussis antigens filamentous hemagglutinin (FHA) and pertactin were observed when BOOSTRIX was administered concomitantly with FLUARIX as compared with BOOSTRIX alone. It is not known if the efficacy of BOOSTRIX is affected by the reduced response to FHA and pertactin.

- When BOOSTRIX is administered concomitantly with other injectable vaccines or Tetanus Immune Globulin, they should be given with separate syringes and at different injection sites. BOOSTRIX should not be mixed with any other vaccine in the same syringe or vial.
- 313 **7.2** Immunosuppressive Therapies
- Immunosuppressive therapies, including irradiation, antimetabolites, alkylating agents,
 cytotoxic drugs, and corticosteroids (used in greater than physiologic doses), may reduce the
 immune response to BOOSTRIX.

317 8 USE IN SPECIFIC POPULATIONS

318 8.1 Pregnancy

319Pregnancy Category C

Animal reproduction studies have not been conducted with BOOSTRIX. It is also not known whether BOOSTRIX can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. BOOSTRIX should be given to a pregnant woman only if clearly needed.

- Animal fertility studies have not been conducted with BOOSTRIX. In a developmental toxicity study, the effect of BOOSTRIX on embryo-fetal and pre-weaning development was evaluated in pregnant rats. Animals were administered INFANRIX prior to gestation and BOOSTRIX during the period of organogenesis (gestation days 6, 8, 11) and later in pregnancy (gestation day 15), 0.1 mL/rat/occasion (a 45-fold increase compared with the human dose of BOOSTRIX on a body weight basis), by intramuscular injection. No adverse effect on pregnancy
- and lactation parameters, embryo-fetal or pre-weaning development was observed. There were

- 331 no fetal malformations or other evidence of teratogenesis noted in this study.
- 332 Pregnancy Registry: GlaxoSmithKline maintains a surveillance registry to collect data
- 333 on pregnancy outcomes and newborn health status outcomes following vaccination with
- 334 BOOSTRIX during pregnancy. Women who receive BOOSTRIX during pregnancy should be
- encouraged to contact GlaxoSmithKline directly or their healthcare provider should contact
- 336 GlaxoSmithKline by calling 1-888-452-9622.

337 8.3 Nursing Mothers

- 338 It is not known whether BOOSTRIX is excreted in human milk. Because many drugs are 339 excreted in human milk, caution should be exercised when BOOSTRIX is administered to a 340 nursing woman.
- 341 8.4 Pediatric Use
- BOOSTRIX is not indicated for use in children younger than 10 years of age. Safety and
 effectiveness of BOOSTRIX in this age group have not been established.

344 8.5 Geriatric Use

345 In clinical trials, 1,104 subjects 65 years of age and older received BOOSTRIX; of these 346 subjects, 299 were 75 years of age and older. In the US elderly (65 years and older) study, 347 immune responses to tetanus and diphtheria toxoids following BOOSTRIX were non-inferior to 348 the comparator Td vaccine. Antibody responses to pertussis antigens following a single dose of 349 BOOSTRIX in the elderly were non-inferior to those observed with INFANRIX administered as 350 a 3-dose series in infants [see Clinical Studies (14.4)]. Solicited adverse events following 351 BOOSTRIX were similar in frequency to those reported with the comparator Td vaccine [see 352 Adverse Reactions (6.1)].

353 11 DESCRIPTION

BOOSTRIX (Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine, Adsorbed) is a noninfectious, sterile, vaccine for intramuscular administration. It contains tetanus toxoid, diphtheria toxoid, and pertussis antigens (inactivated pertussis toxin [PT] and formaldehyde-treated filamentous hemagglutinin [FHA] and pertactin). The antigens are the same as those in INFANRIX, but BOOSTRIX is formulated with reduced quantities of these antigens.

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

The acellular pertussis antigens (PT, FHA, and pertactin) are isolated from *Bordetella pertussis* culture grown in modified Stainer-Scholte liquid medium. PT and FHA are isolated from the fermentation broth; pertactin is extracted from the cells by heat treatment and

- 370 flocculation. The antigens are purified in successive chromatographic and precipitation steps. PT
- is detoxified using glutaraldehyde and formaldehyde. FHA and pertactin are treated withformaldehyde.
- Each antigen is individually adsorbed onto aluminum hydroxide. Each 0.5-mL dose is formulated to contain 5 Lf of tetanus toxoid, 2.5 Lf of diphtheria toxoid, 8 mcg of inactivated DT 8 mag of EUA and 2.5 mag of partectin (60 bile Dalten outer membrane protein)
- PT, 8 mcg of FHA, and 2.5 mcg of pertactin (69 kiloDalton outer membrane protein).
 Tetanus and diphtheria toxoid potency is determined by measuring the amount
- Tetanus and diphtheria toxoid potency is determined by measuring the amount of neutralizing antitoxin in previously immunized guinea pigs. The potency of the acellular pertussis components (inactivated PT and formaldehyde-treated FHA and pertactin) is determined by enzyme-linked immunosorbent assay (ELISA) on sera from previously
- 380 immunized mice.
- Each 0.5-mL dose contains aluminum hydroxide as adjuvant (not more than 0.39 mg aluminum by assay), 4.5 mg of sodium chloride, ≤ 100 mcg of residual formaldehyde, and ≤ 100 mcg of polysorbate 80 (Tween 80).
- BOOSTRIX is available in vials and 2 types of prefilled syringes. One type of prefilled syringe has a tip cap which may contain natural rubber latex and a plunger which does not

synnge has a tip cap which had contain hatting rubber have and a pringer which does not
 contain latex. The other type has a tip cap and a rubber plunger which contain dry natural latex
 rubber. The vial stopper does not contain latex. [See How Supplied/Storage and Handling (16).]

388 12 CLINICAL PHARMACOLOGY

389 **12.1 Mechanism of Action**

- 390 <u>Tetanus:</u> Tetanus is a condition manifested primarily by neuromuscular dysfunction 391 caused by a potent exotoxin released by *C. tetani*. Protection against disease is due to the 392 development of neutralizing antibodies to the tetanus toxin. A serum tetanus antitoxin level of at 393 least 0.01 IU/mL, measured by neutralization assays, is considered the minimum protective 394 level.² A level \ge 0.1 IU/mL by ELISA has been considered as protective.
- 395 <u>Diphtheria:</u> Diphtheria is an acute toxin-mediated infectious disease caused by toxigenic 396 strains of *C. diphtheriae*. Protection against disease is due to the development of neutralizing 397 antibodies to the diphtheria toxin. A serum diphtheria antitoxin level of 0.01 IU/mL, measured 398 by neutralization assays, is the lowest level giving some degree of protection; a level of 399 0.1 IU/mL by ELISA is regarded as protective.³ Diphtheria antitoxin levels \geq 1.0 IU/mL by
- 0.1 IU/mL by ELISA is regarded as protective.⁵ Diphtheria antitoxin levels $\geq 1.0 \text{ IU/mL}$ b 400 ELISA have been associated with long-term protection.³
- 401 <u>Pertussis:</u> Pertussis (whooping cough) is a disease of the respiratory tract caused by 402 *B. pertussis.* The role of the different components produced by *B. pertussis* in either the 403 pertussis of or the immunity to pertussis is not well understand
- 403 pathogenesis of, or the immunity to, pertussis is not well understood.
- 404 13 NONCLINICAL TOXICOLOGY

405 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

BOOSTRIX has not been evaluated for carcinogenic or mutagenic potential, or forimpairment of fertility.

40814CLINICAL STUDIES

The efficacy of the tetanus and diphtheria toxoid components of BOOSTRIX is based on
the immunogenicity of the individual antigens compared to US-licensed vaccines using
established serologic correlates of protection. The efficacy of the pertussis components of
BOOSTRIX was evaluated by comparison of the immune response of adolescents and adults
following a single dose of BOOSTRIX to the immune response of infants following a 3-dose
primary series of INFANRIX. In addition, the ability of BOOSTRIX to induce a booster

415 response to each of the antigens was evaluated.

416 **14.1 Efficacy of INFANRIX**

The efficacy of a 3-dose primary series of INFANRIX in infants has been assessed in 2
clinical studies: A prospective efficacy trial conducted in Germany employing a household
contact study design and a double-blind, randomized, active Diphtheria and Tetanus Toxoids
(DT)-controlled trial conducted in Italy sponsored by the National Institutes of Health (NIH) (for
details see INFANRIX prescribing information). Serological data from a subset of infants

- 422 immunized with INFANRIX in the household contact study were compared with the sera of
- 423 adolescents and adults immunized with BOOSTRIX [see Clinical Studies (14.2, 14.3)]. In the
- 424 household contact study, the protective efficacy of INFANRIX, in infants, against WHO-defined
- 425 pertussis (21 days or more of paroxysmal cough with infection confirmed by culture and/or
- 426 serologic testing) was calculated to be 89% (95% CI: 77%, 95%). When the definition of
- 427 pertussis was expanded to include clinically milder disease, with infection confirmed by culture
- 428 and/or serologic testing, the efficacy of INFANRIX against \geq 7 days of any cough was 67%
- 429 (95% CI: 52%, 78%) and against \geq 7 days of paroxysmal cough was 81% (95% CI: 68%, 89%)
- 430 (for details see INFANRIX prescribing information).

431 **14.2** Immunological Evaluation in Adolescents

432 In a multicenter, randomized, controlled study conducted in the United States, the 433 immune responses to each of the antigens contained in BOOSTRIX were evaluated in sera 434 obtained approximately 1 month after administration of a single dose of vaccine to adolescent 435 subjects (10 to 18 years of age). Of the subjects enrolled in this study, approximately 76% were 436 10 to 14 years of age and 24% were 15 to 18 years of age. Approximately 98% of participants in 437 this study had received the recommended series of 4 or 5 doses of either DTwP or a combination 438 of DTwP and DTaP in childhood. The racial/ethnic demographics were as follows: white 85.8%, 439 black 5.7%, Hispanic 5.6%, Oriental 0.8%, and other 2.1%.

440 <u>Response to Tetanus and Diphtheria Toxoids:</u> The antibody responses to the tetanus
441 and diphtheria toxoids of BOOSTRIX compared with Td vaccine are shown in Table 6. One
442 month after a single dose, anti-tetanus and anti-diphtheria seroprotective rates (≥0.1 IU/mL by
443 ELISA) and booster response rates were comparable between BOOSTRIX and the comparator
444 Td vaccine.

446 Table 6. Antibody Responses to Tetanus and Diphtheria Toxoids Following BOOSTRIX

- 447 Compared With Td Vaccine in Adolescents 10 to 18 Years of Age (ATP Cohort for
- 448 **Immunogenicity**)

				% Booster
		% ≥0.1 IU/mL ^a	% ≥1.0 IU/mL ^a	Response ^b
	Ν	(95% CI)	(95% CI)	(95% CI)
Anti-Tetanus				
BOOSTRIX	2,469-2,516			
Pre-vaccination		97.7 (97.1, 98.3)	36.8 (34.9, 38.7)	—
Post-vaccination		100 (99.8, 100) ^c	99.5 (99.1, 99.7) ^d	89.7 (88.4, 90.8) ^c
Td	817-834			
Pre-vaccination		96.8 (95.4, 97.9)	39.9 (36.5, 43.4)	—
Post-vaccination		100 (99.6, 100)	99.8 (99.1, 100)	92.5 (90.5, 94.2)
Anti-Diphtheria				
BOOSTRIX	2,463-2,515			
Pre-vaccination		85.8 (84.3, 87.1)	17.1 (15.6, 18.6)	—
Post-vaccination		99.9 (99.7, 100) ^c	97.3 (96.6, 97.9) ^d	90.6 (89.4, 91.7) ^c
Td	814-834			
Pre-vaccination		84.8 (82.1, 87.2)	19.5 (16.9, 22.4)	—
Post-vaccination		99.9 (99.3, 100)	99.3 (98.4, 99.7)	95.9 (94.4, 97.2)

449 Td manufactured by MassBioLogics.

- 450 ATP = according-to-protocol; CI = Confidence Interval.
- 451 ^a Measured by ELISA.
- 452 ^b Booster response: In subjects with pre-vaccination <0.1 IU/mL, post-vaccination
- 453 concentration ≥ 0.4 IU/mL. In subjects with pre-vaccination concentration ≥ 0.1 IU/mL, an
- 454 increase of at least 4 times the pre-vaccination concentration.
- 455 ^c Seroprotection rate or booster response rate to BOOSTRIX was non-inferior to Td (upper
 456 limit of two-sided 95% CI on the difference for Td minus BOOSTRIX ≤10%).
- 457 ^d Non-inferiority criteria not prospectively defined for this endpoint.
- 458

459 <u>Response to Pertussis Antigens:</u> The booster response rates of adolescents to the
 460 pertussis antigens are shown in Table 7. For each of the pertussis antigens the lower limit of the
 461 two-sided 95% CI for the percentage of subjects with a booster response exceeded the pre 462 defined lower limit of 80% for demonstration of an acceptable booster response.

		BOOSTRIX
	Ν	% Booster Response ^a (95% CI)
Anti-PT	2,677	84.5 (83.0, 85.9)
Anti-FHA	2,744	95.1 (94.2, 95.9)
Anti-pertactin	2,752	95.4 (94.5, 96.1)

464 Table 7. Booster Responses to the Pertussis Antigens Following BOOSTRIX in Adolescents
465 10 to 18 Years of Age (ATP Cohort for Immunogenicity)

466 ATP = according-to-protocol; CI = Confidence Interval.

467 Booster response: In initially seronegative subjects (<5 EL.U./mL), post-vaccination antibody 468 concentrations ≥ 20 EL.U./mL. In initially seropositive subjects with pre-vaccination antibody 469 concentrations \geq 5 EL.U./mL and \leq 20 EL.U./mL, an increase of at least 4 times the pre-vaccination antibody concentration. In initially seropositive subjects with pre-vaccination 470 471 antibody concentrations ≥20 EL.U./mL, an increase of at least 2 times the pre-vaccination 472 antibody concentration. 473 474 The GMCs to each of the pertussis antigens 1 month following a single dose of BOOSTRIX in the US adolescent study (N = 2,941-2,979) were compared with the GMCs 475 476 observed in infants following a 3-dose primary series of INFANRIX administered at 3, 4, and 477 5 months of age (N = 631-2,884). Table 8 presents the results for the total immunogenicity 478 cohort in both studies (vaccinated subjects with serology data available for at least one pertussis 479 antigen; the majority of subjects in the study of INFANRIX had anti-PT serology data only). 480 These infants were a subset of those who formed the cohort for the German household contact 481 study in which the efficacy of INFANRIX was demonstrated [see Clinical Studies (14.1)]. 482 Although a serologic correlate of protection for pertussis has not been established, anti-PT, anti-483 FHA, and anti-pertactin antibody concentrations observed in adolescents 1 month after a single dose of BOOSTRIX were non-inferior to those observed in infants following a primary 484 485 vaccination series with INFANRIX. 486

487 Table 8. Ratio of GMCs to Pertussis Antigens Following One Dose of BOOSTRIX in

488 Adolescents 10 to 18 Years of Age Compared With 3 Doses of INFANRIX in Infant	s (Total
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489 Immunogenicity Cohort)

	GMC Ratio: BOOSTRIX/INFANRIX	
	(95% CI)	
Anti-PT	$1.90 (1.82, 1.99)^{a}$	
Anti-FHA	7.35 (6.85, 7.89) ^a	
Anti-pertactin	4.19 (3.73, 4.71) ^a	

- GMC = geometric mean antibody concentration, measured in ELISA units; CI = Confidence
 Interval.
- 492 Number of subjects for BOOSTRIX GMC evaluation: Anti-PT = 2,941, anti-FHA = 2,979, and
 493 anti-pertactin = 2,978.
- 494 Number of subjects for INFANRIX GMC evaluation: Anti-PT = 2,884, anti-FHA = 685, and
 495 anti-pertactin = 631.
- ^a GMC following BOOSTRIX was non-inferior to GMC following INFANRIX (lower limit of
 95% CI for the GMC ratio of BOOSTRIX/INFANRIX >0.67).
- 498

499 14.3 Immunological Evaluation in Adults (19 to 64 Years of Age)

500 A multicenter, randomized, observer-blinded study, conducted in the United States, 501 evaluated the immunogenicity of BOOSTRIX compared with the licensed comparator Tdap 502 vaccine (Sanofi Pasteur SA). Vaccines were administered as a single dose to subjects 503 (N = 2,284) who had not received a tetanus-diphtheria booster within 5 years. The immune 504 responses to each of the antigens contained in BOOSTRIX were evaluated in sera obtained 505 approximately 1 month after administration. Approximately 33% of patients were 19 to 29 years 506 of age, 33% were 30 to 49 years of age and 34% were 50 to 64 years of age. Among subjects in the combined vaccine groups, 62% were female; 84% of subjects were white, 8% black, 1% 507 508 Asian, and 7% were of other racial/ethnic groups. 509 Response to Tetanus and Diphtheria Toxoids: The antibody responses to the tetanus 510 and diphtheria toxoids of BOOSTRIX compared with the comparator Tdap vaccine are shown in 511 Table 9. One month after a single dose, anti-tetanus and anti-diphtheria seroprotective rates 512 (≥0.1 IU/mL by ELISA) were comparable between BOOSTRIX and the comparator Tdap

- 513 vaccine.
- 514

515 Table 9. Antibody Responses to Tetanus and Diphtheria Toxoids Following One Dose of

516 **BOOSTRIX** Compared With the Comparator Tdap Vaccine in Adults 19 to 64 Years of

517 Age (ATP Cohort for Immunogenicity)

	N	$\% \ge 0.1 \text{ IU/mL}^{a}$	% ≥1.0 IU/mL ^a
	N	(95% CI)	(95% CI)
Anti-Tetanus			
BOOSTRIX	1,445-1,447		
Pre-vaccination		95.9 (94.8, 96.9)	71.9 (69.5, 74.2)
Post-vaccination		99.6 (99.1, 99.8) ^b	98.3 (97.5, 98.9) ^b
Tdap	727-728		
Pre-vaccination		97.2 (95.8, 98.3)	74.7 (71.4, 77.8)
Post-vaccination		100 (95.5, 100)	99.3 (98.4, 99.8)
Anti-Diphtheria			
BOOSTRIX	1,440-1,444		
Pre-vaccination		85.2 (83.3, 87.0)	23.7 (21.5, 26.0)
Post-vaccination		98.2 (97.4, 98.8) ^b	87.9 (86.1, 89.5) ^c
Tdap	720-727		
Pre-vaccination		89.2 (86.7, 91.3)	26.5 (23.3, 29.9)
Post-vaccination		98.6 (97.5, 99.3)	92.0 (89.8, 93.9)

518 Tdap = Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine, Adsorbed 519 manufactured by Sanofi Pasteur SA.

520 ATP = according-to-protocol; CI = Confidence Interval.

Measured by ELISA. 521 а

b 522 Seroprotection rates for BOOSTRIX were non-inferior to the comparator Tdap vaccine (lower 523 limit of 95% CI on the difference of BOOSTRIX minus Tdap \geq -10%).

524 c Non-inferiority criteria not prospectively defined for this endpoint.

525

526 Response to Pertussis Antigens: Booster response rates to the pertussis antigens are shown in Table 10. For the FHA and pertactin antigens, the lower limit of the 95% CI for the 527 528 booster responses exceeded the pre-defined limit of 80% demonstrating an acceptable booster 529 response following BOOSTRIX. The PT antigen booster response lower limit of the 95% CI 530 (74.9%) did not exceed the pre-defined limit of 80%.

532Table 10. Booster Responses to the Pertussis Antigens Following One Dose of BOOSTRIX

	NI	BOOSTRIX % Booster Response ^a
	N	(95% CI)
Anti-PT	1,419	77.2 (74.9, 79.3) ^b
Anti-FHA	1,433	96.9 (95.8, 97.7) ^c
Anti-pertactin	1,441	93.2 (91.8, 94.4) ^c

533 in Adults 19 to 64 Years of Age (ATP Cohort for Immunogenicity)

534 ATP = according-to-protocol; CI = Confidence Interval.

535 а Booster response: In initially seronegative subjects (<5 EL.U./mL), post-vaccination antibody 536 concentrations ≥ 20 EL.U./mL. In initially seropositive subjects with pre-vaccination antibody 537 concentrations ≥5 EL.U./mL and <20 EL.U./mL, an increase of at least 4 times the pre-538 vaccination antibody concentration. In initially seropositive subjects with pre-vaccination 539 antibody concentrations ≥20 EL.U./mL, an increase of at least 2 times the pre-vaccination 540 antibody concentration. b 541 The PT antigen booster response lower limit of the 95% CI did not exceed the pre-defined 542 limit of 80%. c 543 The FHA and pertactin antigens booster response lower limit of the 95% CI exceeded the pre-544 defined limit of 80%. 545 546 The GMCs to each of the pertussis antigens 1 month following a single dose of 547 BOOSTRIX in the US adult (19 to 64 years of age) study were compared with the GMCs 548 observed in infants following a 3-dose primary series of INFANRIX administered at 3, 4, and 549 5 months of age. Table 11 presents the results for the total immunogenicity cohort in both studies 550 (vaccinated subjects with serology data available for at least one pertussis antigen). These infants 551 were a subset of those who formed the cohort for the German household contact study in which 552 the efficacy of INFANRIX was demonstrated [see Clinical Studies (14.1)]. Although a serologic 553 correlate of protection for pertussis has not been established, anti-PT, anti-FHA, and anti-554 pertactin antibody concentrations observed in adults 1 month after a single dose of BOOSTRIX 555 were non-inferior to those observed in infants following a primary vaccination series with 556 INFANRIX.

- 558 Table 11. Ratio of GMCs to Pertussis Antigens Following One Dose of BOOSTRIX in
- 559 Adults 19 to 64 Years of Age Compared With 3 Doses of INFANRIX in Infants (Total
- 560 Immunogenicity Cohort)

	GMC Ratio: BOOSTRIX/INFANRIX
	(95% CI)
Anti-PT	$1.39(1.32, 1.47)^{a}$
Anti-FHA	7.46 (6.86, 8.12) ^a
Anti-pertactin	3.56 (3.10, 4.08) ^a

- 561 GMC = geometric mean antibody concentration; CI = Confidence Interval.
- Number of subjects for BOOSTRIX GMC evaluation: Anti-PT = 1,460, anti-FHA = 1,472, and
 anti-pertactin = 1,473.
- Number of subjects for INFANRIX GMC evaluation: Anti-PT = 2,884, anti-FHA = 685, and
 anti-pertactin = 631.
- ^a BOOSTRIX was non-inferior to INFANRIX (lower limit of 95% CI for the GMC ratio of
 BOOSTRIX/INFANRIX ≥0.67).
- 568

569 14.4 Immunological Evaluation in the Elderly (65 Years of Age and Older)

570 The US elderly (65 years of age and older) study, a randomized, observer-blinded study, 571 evaluated the immunogenicity of BOOSTRIX (N = 887) compared with a US-licensed 572 comparator Td vaccine (N = 445) (Sanofi Pasteur SA). Vaccines were administered as a single 573 dose to subjects who had not received a tetanus-diphtheria booster within 5 years. Among all 574 vaccine recipients, the mean age was approximately 72 years of age; 54% were female and 95% 575 were white. The immune responses to each of the antigens contained in BOOSTRIX were 576 evaluated in sera obtained approximately 1 month after administration. 577 Response to Tetanus and Diphtheria Toxoids and Pertussis Antigens: Immune 578 responses to tetanus and diphtheria toxoids and pertussis antigens were measured 1 month after 579 administration of a single dose of BOOSTRIX or a comparator Td vaccine. Anti-tetanus and 580 anti-diphtheria seroprotective rates (≥0.1 IU/mL) were comparable between BOOSTRIX and the 581 comparator Td vaccine (Table 12).

583 Table 12. Immune Responses to Tetanus and Diphtheria Toxoids Following BOOSTRIX or

584 Comparator Td Vaccine in the Elderly 65 Years of Age and Older (ATP Cohort for

585 **Immunogenicity**)

	BOOSTRIX	Td
	(N = 844-864)	(N = 430-439)
Anti-T		
% ≥0.1 IU/mL (95% CI)	96.8 (95.4, 97.8) ^a	97.5 (95.6, 98.7)
% ≥1.0 IU/mL (95% CI)	88.8 (86.5, 90.8) ^a	90.0 (86.8, 92.6)
Anti-D		
% ≥0.1 IU/mL (95% CI)	84.9 (82.3, 87.2) ^a	86.6 (83.0, 89.6)
% ≥1.0 IU/mL (95% CI)	52.0 (48.6, 55.4) ^b	51.2 (46.3, 56.0)

Td = Tetanus and Diphtheria Toxoids Adsorbed, a US-licensed Td vaccine, manufactured by
 Sanofi Pasteur SA.

588 ATP = according-to-protocol; CI = Confidence Interval.

^a Seroprotection rates for BOOSTRIX were non-inferior to the comparator Td vaccine (lower
 limit of 95% CI on the difference of BOOSTRIX minus Td ≥-10%).

^b Non-inferiority criteria not prospectively defined for this endpoint.

592

593 The GMCs to each of the pertussis antigens 1 month following a single dose of

594 BOOSTRIX were compared with the GMCs of infants following a 3-dose primary series of

595 INFANRIX administered at 3, 4, and 5 months of age. Table 13 presents the results for the total

596 immunogenicity cohort in both studies (vaccinated subjects with serology data available for at

597 least one pertussis antigen). These infants were a subset of those who formed the cohort for the

598 German household contact study in which the efficacy of INFANRIX was demonstrated *[see*

599 *Clinical Studies (14.1)]*. Although a serologic correlate of protection for pertussis has not been

600 established, anti-PT, anti-FHA, and anti-pertactin antibody concentrations in the elderly

601 (65 years of age and older) 1 month after a single dose of BOOSTRIX were non-inferior to those

602 of infants following a primary vaccination series with INFANRIX.

604 Table 13. Ratio of GMCs to Pertussis Antigens Following One Dose of BOOSTRIX in the

605 Elderly 65 Years of Age and Older Compared With 3 Doses of INFANRIX in Infants

606 (Total Immunogenicity Cohort)

	GMC Ratio: BOOSTRIX/INFANRIX
	(95% CI)
Anti-PT	$1.07 (1.00, 1.15)^{a}$
Anti-FHA	8.24 (7.45, 9.12) ^a
Anti-pertactin	$0.93 (0.79, 1.10)^{a}$

607 GMC = geometric mean antibody concentration; CI = Confidence Interval.

Number of subjects for BOOSTRIX GMC evaluation: Anti-PT = 865, anti-FHA = 847, and anti pertactin = 878.

610 Number of subjects for INFANRIX GMC evaluation: Anti-PT = 2,884, anti-FHA = 685, and 611 anti-pertactin = 631.

^a BOOSTRIX was non-inferior to INFANRIX (lower limit of 95% CI for the GMC ratio of
 BOOSTRIX/INFANRIX ≥0.67).

614

615 14.5 Concomitant Vaccine Administration

616 Concomitant Administration With Meningococcal Conjugate Vaccine: The 617 concomitant use of BOOSTRIX and a tetravalent meningococcal (groups A, C, Y, and W-135) 618 conjugate vaccine (Sanofi Pasteur SA) was evaluated in a randomized study in healthy 619 adolescents 11 to 18 years of age. A total of 1,341 adolescents were vaccinated with 620 BOOSTRIX. Of these, 446 subjects received BOOSTRIX administered concomitantly with 621 meningococcal conjugate vaccine at different injection sites, 446 subjects received BOOSTRIX 622 followed by meningococcal conjugate vaccine 1 month later, and 449 subjects received 623 meningococcal conjugate vaccine followed by BOOSTRIX 1 month later. 624 Immune responses to diphtheria and tetanus toxoids (% of subjects with anti-tetanus and 625 anti-diphtheria antibodies ≥ 1.0 IU/mL by ELISA), pertussis antigens (booster responses and 626 GMCs), and meningococcal antigens (vaccine responses) were measured 1 month (range 30 to 627 48 days) after concomitant or separate administration of BOOSTRIX and meningococcal 628 conjugate vaccine. For BOOSTRIX given concomitantly with meningococcal conjugate vaccine 629 compared to BOOSTRIX administered first, non-inferiority was demonstrated for all antigens, 630 with the exception of the anti-pertactin GMC. The lower limit of the 95% CI for the GMC ratio 631 was 0.54 for anti-pertactin (pre-specified limit \geq 0.67). For the anti-pertactin booster response, 632 non-inferiority was demonstrated. It is not known if the efficacy of BOOSTRIX is affected by 633 the reduced response to pertactin. 634 There was no evidence that BOOSTRIX interfered with the antibody responses to the 635 meningococcal antigens when measured by serum bactericidal assays (rSBA) when given 636 concomitantly or sequentially (meningococcal conjugate vaccine followed by BOOSTRIX or 637 BOOSTRIX followed by meningococcal conjugate vaccine. 638 Concomitant Administration With FLUARIX (Influenza Virus Vaccine): The

- 639 concomitant use of BOOSTRIX and FLUARIX was evaluated in a multicenter, open-label,
- randomized, controlled study of 1,497 adults 19 to 64 years of age. In one group, subjects
- 641 received BOOSTRIX and FLUARIX concurrently (n = 748). The other group received
- 642 FLUARIX at the first visit, then 1 month later received BOOSTRIX (n = 749). Sera was
- obtained prior to and 1 month following concomitant or separate administration of BOOSTRIX
- and/or FLUARIX, as well as 1 month after the separate administration of FLUARIX.
- 645 Immune responses following concurrent administration of BOOSTRIX and FLUARIX
- 646 were non-inferior to separate administration for diphtheria (seroprotection defined as
- $\geq 0.1 \text{ IU/mL}$, tetanus (seroprotection defined as $\geq 0.1 \text{ IU/mL}$ and based on concentrations
- $\geq 1.0 \text{ IU/mL}$, pertussis toxin (PT) antigen (anti-PT GMC) and influenza antigens (percent of
- 649 subjects with hemagglutination-inhibition [HI] antibody titer \geq 1:40 and \geq 4-fold rise in HI titer).
- 650 Non-inferiority criteria were not met for the anti-pertussis antigens FHA and pertactin. The lower
- 651 limit of the 95% CI of the GMC ratio was 0.64 for anti-FHA and 0.60 for anti-pertactin and the
- 652 pre-specified limit was ≥ 0.67 . It is not known if the efficacy of BOOSTRIX is affected by the
- 653 reduced response to FHA and pertactin.

654 **15 REFERENCES**

- Institute of Medicine (IOM). Stratton KR, Howe CJ, Johnston RB, eds. *Adverse events associated with childhood vaccines. Evidence bearing on causality.* Washington, DC:
 National Academy Press; 1994.
- Wassilak SGF, Roper MH, Kretsinger K, and Orenstein WA. Tetanus Toxoid. In: Plotkin
 SA, Orenstein WA, and Offit PA, eds. *Vaccines*. 5th ed. Saunders; 2008:805-839.
- 3. Vitek CR and Wharton M. Diphtheria Toxoid. In: Plotkin SA, Orenstein WA, and Offit PA,
 eds. *Vaccines*. 5th ed. Saunders; 2008:139-156.

662 16 HOW SUPPLIED/STORAGE AND HANDLING

- 663 BOOSTRIX is available in 0.5-mL single-dose vials and disposable prefilled TIP-LOK 664 syringes (packaged without needles):
- 665 NDC 58160-842-01 Vial (contains no latex) in Package of 10: NDC 58160-842-11
- NDC 58160-842-34 Syringe (tip cap may contain latex; plunger contains no latex) in Package of
 1: NDC 58160-842-34
- 668 NDC 58160-842-43 Syringe (tip cap may contain latex; plunger contains no latex) in Package of 669 10: NDC 58160-842-52
- 609 10. NDC 58100-842-52
 - NDC 58160-842-41 Syringe (tip cap and plunger contain latex) in Package of 10: NDC 58160-
 - 671 842-51
 - 672 Store refrigerated between 2° and 8°C (36° and 46°F). Do not freeze. Discard if the
 - 673 vaccine has been frozen.

674 17 PATIENT COUNSELING INFORMATION

- 675 The patient, parent, or guardian should be:
- informed of the potential benefits and risks of immunization with BOOSTRIX.

- 677 informed about the potential for adverse reactions that have been temporally associated with
 678 administration of BOOSTRIX or other vaccines containing similar components.
- instructed to report any adverse events to their healthcare provider.
- informed that safety and efficacy have not been established in pregnant women. Register
- women who receive BOOSTRIX while pregnant in the pregnancy registry by calling 1-888-452-9622.
- e given the Vaccine Information Statements, which are required by the National Childhood
 Vaccine Injury Act of 1986 to be given prior to immunization. These materials are available
- 685 free of charge at the Centers for Disease Control and Prevention (CDC) website
- 686 (www.cdc.gov/vaccines).
- 687
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