Sanofi Pasteur 306 – Adacel[®]

Full Prescribing Information

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

Adacel (Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular Pertussis Vaccine Adsorbed)

Suspension for Intramuscular Injection

Initial US Approval: 2005

-----RECENT MAJOR CHANGES-----

Warnings and Precautions. (5.2)

09/2017

-----DOSAGE AND ADMINISTRATION-----

• A single intramuscular injection of 0.5 mL. (2.1)

-----DOSAGE FORMS AND STRENGTHS-----

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

-----CONTRAINDICATIONS-----

- Severe allergic reaction (eg, anaphylaxis) to any component of Adacel or any other diphtheria toxoid, tetanus toxoid and pertussis antigencontaining vaccine. (4.1)
- Encephalopathy (eg, coma, decreased level of consciousness, prolonged seizures) within 7 days of administration of a previous pertussis antigen-containing vaccine. (4.2)

------WARNINGS AND PRECAUTIONS------

- For one presentation of Adacel, the tip caps of the prefilled syringes may contain natural rubber latex, which may cause allergic reactions in latex sensitive individuals. (5.2, 16)
- If Guillain-Barré syndrome occurred within 6 weeks of receipt of a prior vaccine containing tetanus toxoid, the risk for Guillain-Barré syndrome may be increased following a subsequent dose of Adacel vaccine. (5.3)
- Progressive or unstable neurologic conditions are reasons to defer Adacel vaccination. (5.4)
- Persons who experienced an Arthus-type hypersensitivity reaction following a prior dose of a tetanus toxoid-containing vaccine should not receive Adacel unless at least 10 years have elapsed since the last dose of a tetanus toxoid-containing vaccine. (5.5)

 Syncope (fainting) can occur in association with administration of injectable vaccines, including Adacel. Procedures should be in place to prevent falling injury and manage syncopal reactions.

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

- The most common solicited injection site reactions occurring within 0-14 days following vaccination with Adacel were:
 - For Adolescents 11-17 years of age: pain (77.8%), swelling (20.9%), erythema (20.8%).
 - For Adults 18-64 years of age: pain (65.7%), swelling (21.0%), erythema (24.7%). (6.1)
- The most common solicited systemic reactions occurring within 0-14 days following vaccination with Adacel were:
 - For Adolescents 11-17 years of age: headache (43.7%), body ache or muscle weakness (30.4%), tiredness (15.1%).
 - For Adults 18-64 years of age: headache (33.9%), body ache or muscle weakness (21.9%). (6.1)

- When Adacel vaccine was administered concomitantly with trivalent inactivated influenza vaccine (TIV) to adults 19-64 years of age, a lower antibody response was observed for pertactin antigen as compared to Adacel vaccine administered alone. (7.1, 14.3)
- Immunosuppressive therapies may reduce the immune response to Adacel. (7.2)
- Do not mix Adacel vaccine with any other vaccine in the same syringe or vial.

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

- Safety and effectiveness of Adacel vaccine have not been established in pregnant women. (8.1)
- Pregnancy Surveillance Registry: contact Sanofi Pasteur Inc. at 1-800-822-2463 (1-800-VACCINE). (8.1)

See 17 PATIENT COUNSELING INFORMATION

Revised: [XXX/2017]

FULL PRESCRIBING INFORMATION: CONTENTS*

- 1 INDICATIONS AND USAGE
- 2 DOSAGE AND ADMINISTRATION
 - 2.1 Preparation for Administration
 - 2.2 Administration, Dose and Schedule
 - 2.3 Additional Dosing Information
- 3 DOSAGE FORMS AND STRENGTHS
- 4 CONTRAINDICATIONS
 - 4.1 Hypersensitivity
 - 4.2 Encephalopathy

5 WARNINGS AND PRECAUTIONS

- 5.1 Management of Acute Allergic Reactions
- 5.2 Latex
- 5.3 Guillain-Barré Syndrome and Brachial Neuritis
- 5.4 Progressive or Unstable Neurologic Disorders
- 5.5 Arthus-Type Hypersensitivity
- 5.6 Altered Immunocompetence
- 5.7 Syncope

6 ADVERSE REACTIONS

- 6.1 Clinical Trials Experience
- 6.2 Postmarketing Experience

7 DRUG INTERACTIONS

- 7.1 Concomitant Vaccine Administration
- 7.2 Immunosuppressive Treatments

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

- 8.3 Nursing Mothers
- 8.4 Pediatric Use
- 8.5 Geriatric Use

11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

13 NON-CLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

14 CLINICAL STUDIES

- 14.1 Immunological Evaluation in Adolescents and Adults, 10 Through 64 Years of Age
- 14.2 Concomitant Hepatitis B Vaccine Administration
- 14.3 Concomitant Influenza Vaccine Administration

15 REFERENCES

16 HOW SUPPLIED/STORAGE AND HANDLING

17 PATIENT COUNSELING INFORMATION

* Sections or subsections omitted from the full prescribing information are not listed.

1 FULL PRESCRIBING INFORMATION:

2 1 INDICATIONS AND USAGE

- 3 Adacel® is a vaccine indicated for active booster immunization against tetanus, diphtheria and
- 4 pertussis. Adacel vaccine is approved for use as a single dose in individuals 10 through 64 years
- 5 of age.

6 2 DOSAGE AND ADMINISTRATION

7 **2.1 Preparation for Administration**

- 8 Just before use, shake the vial or syringe well until a uniform, white, cloudy suspension results.
- 9 Parenteral drug products should be inspected visually for particulate matter and discoloration
- prior to administration, whenever solution and container permit. If either of these conditions exist,
- 11 the vaccine should not be administered.
- When withdrawing a dose from a stoppered vial, do not remove either the stopper or the metal
- seal holding it in place. Use a separate sterile needle and syringe for each injection. Using a sterile
- 14 needle and syringe, withdraw the 0.5 mL dose of vaccine from the single-dose vial and administer
- 15 the vaccine to the individual. Changing needles between withdrawing the vaccine from the vial
- and injecting it into a recipient is not necessary unless the needle has been damaged or
- 17 contaminated.
- Adacel vaccine should not be combined through reconstitution or mixed with any other vaccine.

19 **2.2** Administration, Dose and Schedule

- Adacel vaccine is administered as a single 0.5 mL intramuscular injection into the deltoid muscle
- 21 of the upper arm.
- 22 Do not administer this product intravenously, subcutaneously or intradermally.
- 23 There are no data to support repeat administration of Adacel vaccine.
- 24 Five years should have elapsed since the recipient's last dose of tetanus toxoid, diphtheria toxoid
- and/or pertussis containing vaccine and the administration of Adacel vaccine.

27 **2.3 Additional Dosing Information**

- 28 **Primary series:** The safety and effectiveness of Adacel vaccine used as a primary series or to
- 29 complete the primary series, for diphtheria, tetanus, or pertussis has not been demonstrated.
- 30 Wound management: If tetanus prophylaxis is needed for wound management, Adacel may be
- 31 given if no previous dose of any Tetanus Toxoid, Reduced Diphtheria Toxoid and Acellular
- 32 Pertussis Vaccine, Adsorbed (Tdap) has been administered.

33 3 DOSAGE FORMS AND STRENGTHS

- Adacel vaccine is a suspension for injection (0.5 mL dose) available in 0.5 mL single-dose vials
- 35 and prefilled syringes. [See DOSAGE AND ADMINISTRATION (2.2) and HOW
- 36 SUPPLIED/STORAGE AND HANDLING (16).]

37 4 CONTRAINDICATIONS

38 4.1 Hypersensitivity

- 39 A severe allergic reaction (eg, anaphylaxis) after a previous dose of any tetanus toxoid, diphtheria
- 40 toxoid or pertussis containing vaccine or any other component of this vaccine is a contraindication
- 41 to administration of Adacel vaccine. [See DESCRIPTION (11).] Because of uncertainty as to
- 42 which component of the vaccine may be responsible, none of the components should be
- 43 administered. Alternatively, such individuals may be referred to an allergist for evaluation if
- 44 further immunizations are to be considered.

45 **4.2 Encephalopathy**

- Encephalopathy (eg, coma, prolonged seizures, or decreased level of consciousness) within 7 days
- of a previous dose of a pertussis containing vaccine not attributable to another identifiable cause is
- 48 a contraindication to administration of any pertussis containing vaccine, including
- 49 Adacel vaccine.

51

50 5 WARNINGS AND PRECAUTIONS

5.1 Management of Acute Allergic Reactions

- 52 Epinephrine hydrochloride solution (1:1,000) and other appropriate agents and equipment must be
- available for immediate use in case an anaphylactic or acute hypersensitivity reaction occurs.
- **54 5.2 Latex**
- For one presentation of Adacel, the tip caps of the prefilled syringes may contain natural rubber
- latex, which may cause allergic reactions in latex sensitive individuals. The vial stopper is not
- 57 made with natural rubber latex. [See HOW SUPPLIED/STORAGE AND HANDLING (16).]

58 **5.3 Guillain-Barré Syndrome and Brachial Neuritis**

- A review by the Institute of Medicine found evidence for acceptance of a causal relation between
- 60 tetanus toxoid and both brachial neuritis and Guillain-Barré syndrome. (1) If Guillain-Barré
- 61 syndrome occurred within 6 weeks of receipt of prior vaccine containing tetanus toxoid, the
- 62 risk for Guillain-Barré syndrome may be increased following a dose of Adacel vaccine.

63 5.4 Progressive or Unstable Neurologic Disorders

- Progressive or unstable neurologic conditions are reasons to defer Adacel. It is not known whether
- administration of Adacel to persons with an unstable or progressive neurologic disorder might
- hasten manifestations of the disorder or affect the prognosis. Administration of Adacel to persons
- with an unstable or progressive neurologic disorder may result in diagnostic confusion between
- 68 manifestations of the underlying illness and possible adverse effects of vaccination.

69 5.5 Arthus-Type Hypersensitivity

- 70 Persons who experienced an Arthus-type hypersensitivity reaction following a prior dose of a
- 71 tetanus toxoid-containing vaccine should not receive Adacel unless at least 10 years have elapsed
- since the last dose of a tetanus toxoid containing vaccine.

73 **5.6 Altered Immunocompetence**

- 74 If Adacel vaccine is administered to immunocompromised persons, including persons receiving
- 75 immunosuppressive therapy, the expected immune response may not be obtained. [See *Drug*
- 76 *Interactions* (7.2).]

5.7 Syncope

77

81

- 78 Syncope (fainting) can occur in association with administration of injectable vaccine, including
- Adacel. Procedures should be in place to prevent falling injury and manage syncopal reactions.

80 6 ADVERSE REACTIONS

6.1 Clinical Trials Experience

- 82 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
- 84 of another vaccine and may not reflect the rates observed in practice. The adverse reaction
- 85 information from clinical trials does, however, provide a basis for identifying the adverse events
- 86 that appear to be related to vaccine use and for approximating rates of those events. As with any
- vaccine, there is the possibility that broad use of Adacel vaccine could reveal adverse reactions
- 88 not observed in clinical trials.
- The safety of Adacel vaccine was evaluated in 5 clinical studies. A total of 7,143 individuals 10
- 90 through 64 years of age inclusive (4,695 adolescents 10 through 17 years of age and, 2,448 adults
- 91 18 through 64 years of age) received a single dose of Adacel vaccine.
- 92 Clinical study Td506 was a randomized, observer-blind, active controlled trial that enrolled
- adolescents 11 through 17 years of age (Adacel vaccine N = 1,184; Td vaccine N = 792) and
- adults 18 through 64 years of age (Adacel vaccine N = 1,752; Td vaccine N = 573). Study
- participants had not received tetanus or diphtheria containing vaccines within the previous 5
- 96 years. Solicited local and systemic reactions and unsolicited adverse events were monitored daily
- 97 for 14 days post-vaccination using a diary card. From days 14-28 post-vaccination, information on
- 98 adverse events necessitating a medical contact, such as a telephone call, visit to an emergency
- 99 room, physician's office or hospitalization, was obtained via telephone interview or at an interim
- clinic visit. From days 28 to 6 months post-vaccination, participants were monitored for
- unexpected visits to a physician's office or to an emergency room, onset of serious illness and
- hospitalizations. Information regarding adverse events that occurred in the 6 month post-
- vaccination time period was obtained from participants via telephone contact. At least 96% of
- participants completed the 6-month follow-up evaluation.

Solicited Adverse Events in the US Adolescent and Adult Study (Td506)

The frequency of selected solicited adverse events (erythema, swelling, pain and fever) occurring during days 0-14 following vaccination with Adacel vaccine or Td vaccine in adolescents 11 through 17 years of age and adults 18 through 64 years of age are presented in Table 1. Most of these events were reported at a similar frequency in recipients of both Adacel vaccine and Td vaccine. Pain at the injection site was the most common adverse reaction in 62.9% to 77.8% of all vaccinees. In addition, overall rates of pain were higher in adolescent recipients of Adacel vaccine compared to Td vaccine recipients. Rates of moderate and severe pain in adolescents did not significantly differ between the Adacel vaccine and Td vaccine groups. Among adults the rates of pain, after receipt of Adacel vaccine or Td vaccine, did not significantly differ. Fever of 38°C and higher was uncommon, although in the adolescent age group, it occurred significantly more frequently in Adacel vaccine recipients than Td vaccine recipients.

Table 1: Frequencies of Solicited Injection Site Reactions and Fever for Adolescents and Adults, Days 0-14, Following Vaccination With Adacel Vaccine or Td Vaccine in Study Td506

		Adolescents		Adults		
		11-17 years		18-64 y	ears	
		Adacel	\mathbf{Td}^{\ddagger}	Adacel	\mathbf{Td}^{\ddagger}	
		$N^{\dagger} = 1,170-1,175$	$\mathbf{N}^{\dagger} = 783 - 787$	$N^{\dagger} = 1,688-1,698$	$N^{\dagger} = 551-561$	
A	dverse Event*	(%)	(%)	(%)	(%)	
Injection	Any	77.8 [§]	71.0	65.7	62.9	
Site	Moderate**	18.0	15.6	15.1	10.2	
Pain	Severe ^{††}	1.5	0.6	1.1	0.9	
	Any	20.9	18.3	21.0	17.3	
	Moderate**			•		
Injection Site	1.0 to 3.4 cm	6.5	5.7	7.6	5.4	
Swelling	Severe ^{††}		1	1		
8	≥3.5 cm	6.4	5.5	5.8	5.5	
	≥5 cm (2 inches)	2.8	3.6	3.2	2.7	
	Any	20.8	19.7	24.7	21.6	
T	Moderate**					
Injection Site	1.0 to 3.4 cm	5.9	4.6	8.0	8.4	
Erythema	Severe ^{††}			•		
	≥3.5 cm	6.0	5.3	6.2	4.8	
	≥5 cm (2 inches)	2.7	2.9	4.0	3.0	
	≥38.0°C (≥100.4°F)	5.0 [§]	2.7	1.4	1.1	
Fever	≥38.8°C to ≤39.4°C	0.9	0.6	0.4	0.2	
	(≥102.0°F to ≤103.0°F)		0.0		0.2	
	≥39.5°C (≥103.1°F)	0.2	0.1	0.0	0.2	

^{*} The study sample size was designed to detect >10% differences between Adacel and Td vaccines for events of 'Any' intensity.

 $^{^{\}dagger}$ N = number of participants with available data.

[‡] Tetanus and Diphtheria Toxoids Adsorbed for Adult Use manufactured by Sanofi Pasteur Inc., Swiftwater, PA.

- Adacel vaccine did not meet the non-inferiority criterion for rates of 'Any' Pain in adolescents compared to Td vaccine rates (upper limit of the 95% CI on the difference for Adacel vaccine minus Td vaccine was 10.7% whereas the criterion was <10%). For 'Any' Fever the non-inferiority criteria was met, however, 'Any' Fever was statistically higher in adolescents receiving Adacel vaccine.
- Interfered with activities, but did not necessitate medical care or absenteeism.
- Incapacitating, prevented the performance of usual activities, may have/or did necessitate medical care or absenteeism.
- The frequency of other solicited adverse events (days 0-14) are presented in Table 2. The rates of
- these events following Adacel vaccine were comparable with those observed with Td vaccine.
- Headache was the most frequent systemic reaction and was usually of mild to moderate intensity.

Table 2: Frequencies of Other Solicited Adverse Events for Adolescents and Adults, Days 0 14, Following Vaccination With Adacel Vaccine or Td Vaccine in Study Td506

Adverse Event		Adolescents 1	1-17 years	Adults 18-64 years		
		Adacel	\mathbf{Td}^{\dagger}	Adacel	\mathbf{Td}^{\dagger}	
		$N^* = 1,174-1,175$	$N^* = 787$	$N^* = 1,697-1,698$		
		(%)	(%)	(%)	(%)	
	Any	43.7	40.4	33.9	34.1	
Headache	Moderate [‡]	14.2	11.1	11.4	10.5	
	Severe [§]	2.0	1.5	2.8	2.1	
Body Ache	Any	30.4	29.9	21.9	18.8	
or Muscle	Moderate [‡]	8.5	6.9	6.1	5.7	
Weakness	Severe [§]	1.3	0.9	1.2	0.9	
	Any	30.2	27.3	24.3	20.7	
Tiredness	Moderate [‡]	9.8	7.5	6.9	6.1	
	Severe [§]	1.2	1.0	1.3	0.5	
	Any	15.1	12.6	8.1	6.6	
Chills	Moderate [‡]	3.2	2.5	1.3	1.6	
	Seve re [§]	0.5	0.1	0.7	0.5	
Sore and	Any	11.3	11.7	9.1	7.0	
Swollen	Moderate [‡]	2.6	2.5	2.5	2.1	
Joints	Severe [§]	0.3	0.1	0.5	0.5	
	Any	13.3	12.3	9.2	7.9	
Nausea	Moderate [‡]	3.2	3.2	2.5	1.8	
	Seve re [§]	1.0	0.6	0.8	0.5	
Lymph	Any	6.6	5.3	6.5	4.1	
Node	Moderate [‡]	1.0	0.5	1.2	0.5	
Swelling	Severe [§]	0.1	0.0	0.1	0.0	
	Any	10.3	10.2	10.3	11.3	
Diarrhea	Moderate [‡]	1.9	2.0	2.2	2.7	
	Seve re [§]	0.3	0.0	0.5	0.5	
	Any	4.6	2.8	3.0	1.8	
Vomiting	Moderate [‡]	1.2	1.1	1.0	0.9	
	Severe [§]	0.5	0.3	0.5	0.2	
Rash	Any	2.7	2.0	2.0	2.3	

N = number of participants with available data.

[†] Tetanus and Diphtheria Toxoids Adsorbed for Adult Use manufactured by Sanofi Pasteur Inc., Swiftwater, PA.

[‡] Interfered with activities, but did not necessitate medical care or absenteeism.

[§] Incapacitating, prevented the performance of usual activities, may have/or did necessitate medical care or absenteeism.

Injection site and systemic solicited reactions occurred at similar rates in Adacel vaccine and
Td vaccine recipients in the 3 day post-vaccination period. Most injection site reactions occurred
within the first 3 days after vaccination (with a mean duration of less than 3 days). The rates of
unsolicited adverse events reported from days 14-28 post-vaccination were comparable between
the two vaccine groups, as were the rates of unsolicited adverse events from day 28 through 6
months. There were no spontaneous reports of extensive limb swelling of the injected limb in
study Td506, nor in the other three studies which also contributed to the safety database for
Adacel vaccine.
Injection Site and Systemic Reactions When Given With Hepatitis B Vaccine
In the concomitant vaccination study with Adacel and Hepatitis B vaccines [see Clinical
Studies (14)], injection site and systemic adverse events were monitored daily for 14 days post-
vaccination using a diary card. Injection site adverse events were only monitored at site/arm of
Adacel vaccine administration. Unsolicited reactions (including immediate reactions, serious
adverse events and events that elicited seeking medical attention) were collected at a clinic visit or
via telephone interview for the duration of the trial, ie, up to 6 months post-vaccination.
The rates reported for fever and injection site pain (at the Adacel vaccine administration site) were
similar when Adacel and Hep B vaccines were given concurrently or separately. However, the
rates of injection site erythema (23.4% for concomitant vaccination and 21.4% for separate
administration) and swelling (23.9% for concomitant vaccination and 17.9% for separate
administration) at the Adacel vaccine administration site were increased when co-administered.
Swollen and/or sore joints were reported by 22.5% for concomitant vaccination and 17.9% for
separate administration. The rates of generalized body aches in the individuals who reported
swollen and/or sore joints were 86.7% for concomitant vaccination and 72.2% for separate
administration. Most joint complaints were mild in intensity with a mean duration of 1.8 days.
The incidence of other solicited and unsolicited adverse events were not different between the
2 study groups.
Injection Site and Systemic Reactions When Given With Trivalent Inactivated Influenza
Vaccine (TIV)
In the concomitant vaccination study with Adacel vaccine and trivalent inactivated influenza
vaccine [see Clinical Studies (14)], injection site and systemic adverse events were monitored for

155 14 days post-vaccination using a diary card. All unsolicited reactions occurring through day 14 156 were collected. From day 14 to the end of the trial, ie, up to 84 days, only events that elicited 157 seeking medical attention were collected. 158 The rates of fever and injection site erythema and swelling were similar for recipients of 159 concurrent and separate administration of Adacel vaccine and TIV. However, pain at the Adacel 160 vaccine injection site occurred at statistically higher rates following concurrent administration 161 (66.6%) versus separate administration (60.8%). The rates of sore and/or swollen joints were 162 13% for concurrent administration and 9% for separate administration. Most joint complaints 163 were mild in intensity with a mean duration of 2.0 days. The incidence of other solicited and 164 unsolicited adverse events were similar between the 2 study groups. 165 **Additional Studies** 166 In an additional study, 1,806 adolescents 11 through 17 years of age received Adacel vaccine as 167 part of the lot consistency study used to support Adacel vaccine licensure. This study was a 168 randomized, double-blind, multi-center trial designed to assess lot consistency as measured by the 169 safety and immunogenicity of 3 lots of Adacel vaccine when given as a booster dose to 170 adolescents 11 through 17 years of age inclusive. Local and systemic adverse events were 171 monitored for 14 days post-vaccination using a diary card. Unsolicited adverse events and serious 172 adverse events were collected for 28 days post-vaccination. Pain was the most frequently reported 173 local adverse event occurring in approximately 80% of all participants. Headache was the most 174 frequently reported systemic event occurring in approximately 44% of all participants. Sore 175 and/or swollen joints were reported by approximately 14% of participants. Most joint complaints 176 were mild in intensity with a mean duration of 2.0 days. 177 An additional 962 adolescents and adults received Adacel vaccine in three supportive Canadian 178 studies used as the basis for licensure in other countries. Within these clinical trials, the rates of 179 local and systemic reactions following Adacel vaccine were similar to those reported in the four 180 principal trials in the US with the exception of a higher rate (86%) of adults experiencing 'any' 181 local injection site pain. The rate of severe pain (0.8%), however, was comparable to the rates 182 reported in four principal trials conducted in the US. There was one spontaneous report of whole-183 arm swelling of the injected limb among the 277 Td vaccine recipients, and two spontaneous 184 reports among the 962 Adacel vaccine recipients in the supportive Canadian studies.

185	An additional study, Td519, enrolled 1,302 individuals in an open label, two-arm, multi-center
186	trial (651 subjects in each group) to evaluate the safety and immunogenicity of a single dose of
187	Adacel administered to persons 10 to <11 years of age compared to persons 11 to <12 years of
188	age. Immediate reactions were monitored for 20 minutes post-vaccination. Solicited local and
189	systemic adverse events were monitored for 7 days post-vaccination using a diary card.
190	Unsolicited and serious adverse events were collected for approximately 30 days post-
191	vaccination. Similar rates of immediate, solicited and unsolicited adverse reactions were reported
192	in each of the two age cohorts. One serious adverse event, not related to vaccination, was reported
193	in the younger age group.
194	Serious Adverse Events in All Safety Studies
195	In all the studies, participants were monitored for serious adverse events throughout the duration
196	of the study.
197	Throughout the 6-month follow-up period in study Td506, serious adverse events were reported in
198	1.5% of Adacel vaccine recipients and in 1.4% of Td vaccine recipients. Two serious adverse
199	events in adults were neuropathic events that occurred within 28 days of Adacel vaccine
200	administration; one severe migraine with unilateral facial paralysis and one diagnosis of nerve
201	compression in neck and left arm. Similar or lower rates of serious adverse events were reported
202	in the other trials in participants up to 64 years of age and no additional neuropathic events were
203	reported.
204	6.2 Postmarketing Experience
205	The following adverse events of Adacel have been spontaneously reported in the US and other
206	countries. Because these events are reported voluntarily from a population of uncertain size, it
207	may not be possible to reliably estimate their frequency or establish a causal relationship to
208	vaccine exposure.
209	The following adverse events were included based on one or more of the following factors:
210	severity, frequency of reporting or strength of evidence for a causal relationship to Adacel
211	vaccine.
212	Immune system disorders
213	Anaphylactic reaction, hypersensitivity reaction (angioedema, edema, rash, hypotension)

214	Nervous system disorders	
215	Paraesthesia, hypoesthesia, Guillain-Barré syndrome, brachial neuritis, facial palsy,	
216	convulsion, syncope, myelitis	
217	Cardiac disorders	
218	Myocarditis	
219	Skin and subcutaneous tissue disorders	
220	Pruritus, urticaria	
221	Musculoskeletal and connective tissue disorders	
222	Myositis, muscle spasm	
223	General disorders and administration site conditions	
224	Large injection site reactions (>50 mm), extensive limb swelling from the injection site	
225	beyond one or both joints	
226	Injection site bruising, sterile abscess	
227	7 DRUG INTERACTIONS	
228	7.1 Concomitant Vaccine Administration	
229	When Adacel vaccine is administered concomitantly with other injectable vaccines or Tetanus	
230	Immune Globulin, they should be given with separate syringes and at different injection sites.	
231	Adacel should not be mixed with any other vaccine in the same syringe or vial.	
232	In clinical studies, Adacel vaccine was administered concomitantly with one of the following US	\ -
233	licensed vaccines: Hepatitis B (10 mcg, two dose regimen) or trivalent inactivated influenza	
234	vaccines (TIV). [See Adverse Reactions (6.1) and Clinical Studies (14).]	
235	Hepatitis B Vaccine	
236	Concomitant immunization of Adacel vaccine with Hepatitis B vaccine did not result in reduced	
237	antibody responses to any of the antigens from either vaccine.	

Trivalent Inactivated Influenza Vaccine (TIV)
No interference in tetanus and diphtheria seroprotection rates and responses to influenza vaccine,
detoxified pertussis toxin (PT), fimbriae types 2 and 3 (FIM) or filamentous hemagglutinin (FHA)
were observed when Adacel vaccine was administered concomitantly with TIV compared to
separate administration. A lower pertactin (PRN) GMC was observed when Adacel vaccine was
administered concomitantly with TIV compared to separate administration.
7.2 Immunosuppressive Treatments
Immunosuppressive therapies, including irradiation, antimetabolites, alkylating agents, cytotoxic
drugs and corticosteroids (used in greater than physiologic doses), may reduce the immune
response to vaccines. [See Warnings and Precautions (5.6).]

273

8 USE IN SPECIFIC POPULATIONS

250	8.1 Pregnancy
251	Pregnancy Category C
252	Animal reproduction studies have not been conducted with Adacel vaccine. It is also not known
253	whether Adacel vaccine can cause fetal harm when administered to a pregnant woman or can
254	affect reproduction capacity. Adacel vaccine should be given to a pregnant woman only if clearly
255	needed.
256	Animal fertility studies have not been conducted with Adacel vaccine. The effect of Adacel
257	vaccine on embryo-fetal and pre-weaning development was evaluated in two developmental
258	toxicity studies using pregnant rabbits. Animals were administered Adacel vaccine twice prior to
259	gestation, during the period of organogenesis (gestation day 6) and later during pregnancy on
260	gestation day 29, 0.5 mL/rabbit/occasion (a 17-fold increase compared to the human dose of
261	Adacel vaccine on a body weight basis), by intramuscular injection. No adverse effects on
262	pregnancy, parturition, lactation, embryo-fetal or pre-weaning development were observed. There
263	were no vaccine related fetal malformations or other evidence of teratogenesis noted in this study.
264	Registry of Receipt of Adacel Vaccine During Pregnancy
265	Sanofi Pasteur Inc. maintains a surveillance registry to collect data on pregnancy outcomes and
266	newborn health status outcomes following vaccination with Adacel vaccine during pregnancy.
267	Women who receive Adacel vaccine during pregnancy are encouraged to contact directly or have
268	their health-care professional contact Sanofi Pasteur Inc. at 1-800-822-2463 (1-800-VACCINE).
269	8.3 Nursing Mothers
270	It is not known whether Adacel vaccine is excreted in human milk. Because many drugs are
271	excreted in human milk, caution should be exercised when Adacel vaccine is given to a nursing
272	woman.

285

8.4 Pediatric Use

- 275 Adacel vaccine is not approved for individuals less than 10 years of age. Safety and effectiveness
- of Adacel vaccine in persons less than 10 years of age have not been established.

277 8.5 Geriatric Use

- Adacel vaccine is not approved for use in individuals 65 years of age and older.
- 279 In a clinical study, individuals 65 years of age and older received a single dose of Adacel vaccine.
- 280 Based on pre-specified criteria, persons 65 years of age and older who received a dose of Adacel
- vaccine had lower geometric mean concentrations of antibodies to PT, PRN and FIM when
- compared to infants who had received a primary series of DAPTACEL®, Diphtheria and Tetanus
- 283 Toxoids and Acellular Pertussis Vaccine Adsorbed (DTaP). [See Section 14 for description of
- 284 DAPTACEL vaccine.]

11 DESCRIPTION

- Adacel vaccine is a sterile isotonic suspension of tetanus and diphtheria toxoids and pertussis
- antigens adsorbed on aluminum phosphate, for intramuscular injection.
- Each 0.5 mL dose contains 5 Lf tetanus toxoid (T), 2 Lf diphtheria toxoid (d), and acellular
- pertussis antigens [2.5 mcg detoxified pertussis toxin (PT), 5 mcg filamentous hemagglutinin
- 290 (FHA), 3 mcg pertactin (PRN), 5 mcg fimbriae types 2 and 3 (FIM)]. Other ingredients per 0.5
- 291 mL dose include 1.5 mg aluminum phosphate (0.33 mg aluminum) as the adjuvant, ≤5 mcg
- residual formaldehyde, <50 ng residual glutaraldehyde and 3.3 mg (0.6% v/v) 2-phenoxyethanol
- 293 (not as a preservative). The antigens are the same as those in DAPTACEL vaccine; however,
- 294 Adacel vaccine is formulated with reduced quantities of diphtheria and detoxified PT.
- 295 The acellular pertussis vaccine components are produced from *Bordetella pertussis* cultures
- 296 grown in Stainer-Scholte medium (2) modified by the addition of casamino acids and dimethyl-
- beta-cyclodextrin. PT, FHA and PRN are isolated separately from the supernatant culture
- 298 medium. FIM are extracted and co-purified from the bacterial cells. The pertussis antigens are
- 299 purified by sequential filtration, salt-precipitation, ultrafiltration and chromatography. PT is
- detoxified with glutaraldehyde, FHA is treated with formaldehyde, and the residual aldehydes are
- removed by ultrafiltration. The individual antigens are adsorbed onto aluminum phosphate.
- 302 The tetanus toxin is produced from *Clostridium tetani* grown in modified Mueller-Miller

303	casamino acid medium without beef heart infusion. (3) Tetanus toxin is detoxified with
304	$formal dehyde \ and \ purified \ by \ ammonium \ sulfate \ fractionation \ and \ diafiltration. \ {\it Corynebacterium}$
305	diphtheriae is grown in modified Mueller's growth medium. (4) After purification by ammonium
306	sulfate fractionation, diphtheria toxin is detoxified with formaldehyde and diafiltered.
307	The adsorbed diphtheria, tetanus and acellular pertussis components are combined with aluminum
308	phosphate (as adjuvant), 2-phenoxyethanol (not as a preservative) and water for injection. Adacel
309	vaccine does not contain a preservative.
310	In the guinea pig potency test, the tetanus component induces at least 2 neutralizing units/mL of
311	serum and the diphtheria component induces at least 0.5 neutralizing units/mL of serum. The
312	potency of the acellular pertussis vaccine components is evaluated by the antibody response of
313	immunized mice to detoxified PT, FHA, PRN and FIM as measured by enzyme-linked
314	immunosorbent assay (ELISA).
315	Diphtheria and tetanus toxoids are individually adsorbed onto aluminum phosphate.
316	

339

fertility.

317	12 CLINICAL PHARMACOLOGY
318	12.1 Mechanism of Action
319	Tetanus
320	Tetanus is a disease manifested primarily by neuromuscular dysfunction caused by a potent
321	exotoxin released by C tetani.
322	Protection against disease is due to the development of neutralizing antibodies to tetanus toxin. A
323	serum tetanus antitoxin level of at least 0.01 IU/mL, measured by neutralization assay is
324	considered the minimum protective level. (5) (6)
325	Diphtheria
326	Diphtheria is an acute toxin-mediated disease caused by toxigenic strains of C diphtheriae.
327	Protection against disease is due to the development of neutralizing antibodies to diphtheria toxin.
328	A serum diphtheria antitoxin level of 0.01 IU/mL is the lowest level giving some degree of
329	protection. Antitoxin levels of at least 0.1 IU/mL are generally regarded as protective. (5) Levels
330	of 1.0 IU/mL have been associated with long-term protection. (7)
331	Pertussis
332	Pertussis (whooping cough) is a respiratory disease caused by <i>B pertussis</i> . This Gram-negative
333	coccobacillus produces a variety of biologically active components, though their role in either the
334	pathogenesis of, or immunity to, pertussis has not been clearly defined.
335	13 NON-CLINICAL TOXICOLOGY
336	13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
337	Adacel vaccine has not been evaluated for carcinogenic or mutagenic potential, or impairment of

340	14 CLINICAL STUDIES
341	The efficacy of the tetanus toxoid and diphtheria toxoid used in Adacel vaccine was based on the
342	immune response to these antigens compared to a US licensed Tetanus and Diphtheria Toxoids
343	Adsorbed For Adult Use (Td) vaccine manufactured by Sanofi Pasteur Inc., Swiftwater, PA. The
344	primary measures for immune response to the diphtheria and tetanus toxoids were the percentage
345	of participants attaining an antibody level of at least 0.1 IU/mL.
346	The efficacy of the pertussis antigens used in Adacel vaccine was inferred based on a comparison
347	of pertussis antibody levels achieved in recipients of a single booster dose of Adacel vaccine with
348	those obtained in infants after three doses of DAPTACEL vaccine. In the Sweden I Efficacy Trial
349	three doses of DAPTACEL vaccine were shown to confer a protective efficacy of 84.9% (95%
350	CI: 80.1%, 88.6%) against WHO defined pertussis (21 days of paroxysmal cough with laboratory-
351	confirmed B pertussis infection or epidemiological link to a confirmed case). The protective
352	efficacy against mild pertussis (defined as at least one day of cough with laboratory-confirmed
353	B pertussis infection) was 77.9% (95% CI: 72.6%, 82.2%). (8)
354	In addition, the ability of Adacel vaccine to elicit a booster response (defined as rise in antibody
355	concentration after vaccination) to the tetanus, diphtheria and pertussis antigens following
356	vaccination was evaluated. The demonstration of a booster response depended on the antibody
357	concentration to each antigen as established based on the 95th percentile of the pre-vaccination
358	antibody concentrations observed in historical clinical trials with Adacel vaccine.
359	14.1 Immunological Evaluation in Adolescents and Adults, 10 Through 64 Years of
360	Age
361	Study Td506 was a comparative, multi-center, randomized, observer-blind, controlled trial which
362	enrolled 4,480 participants; 2,053 adolescents (11 through 17 years of age) and 2,427 adults (18
363	through 64 years of age). Enrollment was stratified by age to ensure adequate representation
364	across the entire age range. Participants had not received a tetanus or diphtheria toxoid containing
365	vaccine within the previous 5 years. After enrollment participants were randomized to receive one
366	dose of either Adacel vaccine or Td vaccine. A total of 4,461 randomized participants were
367	vaccinated. The per-protocol immunogenicity subset included 1,270 Adacel vaccine recipients
368	and 1,026 Td vaccine recipients. Sera were obtained before and approximately 35 days after

369	vaccination. [Blinding procedures for safety assessments are described in ADVERSE REACTIONS
370	(6).]
371	Demographic characteristics were similar within age groups and between the vaccine groups. A
372	total of 76% of the adolescents and 1.1% of the adults reported a history of receiving 5 previous
373	doses of diphtheria-tetanus-pertussis containing vaccines. Anti-tetanus and anti-diphtheria
374	seroprotection rates (\geq 0.1 IU/mL) and booster response rates were comparable between Adacel
375	and Td vaccines. (See Table 3 and Table 4.) Adacel vaccine induced pertussis antibody levels that
376	were non-inferior to those of Swedish infants who received three doses of DAPTACEL vaccine.
377	(See Table 5.) Acceptable booster responses to each of the pertussis antigens were also
378	demonstrated, ie, the percentage of participants with a booster response exceeded the pre-defined
379	lower limit. (See Table 6.)

- Table 3: Pre-vaccination and Post-vaccination Antibody Responses and Booster Response
- 381 Rates to Tetanus Toxoid Following Adacel Vaccine as Compared to Td Vaccine in
- 382 Adolescents and Adults 11 Through 64 Years of Age

			Tetanus Antitoxin (IU/mL)				
			Pre-vaccination		1 Month Post-vaccination		ation
Age Group (years)	Vaccine	N*	% ≥0.10 (95% CI)	% ≥1.0 (95% CI)	% ≥0.10 (95% CI)	% ≥1.0 (95% CI)	% Booster [†] (95% CI)
11-17	Adacel	527	99.6 (98.6, 100.0)	44.6 (40.3, 49.0)	100.0 [‡] (99.3, 100.0)	99.6 [§] (98.6, 100.0)	91.7 [‡] (89.0, 93.9)
	Td**	516	99.2 (98.0, 99.8)	43.8 (39.5, 48.2)	100.0 (99.3, 100.0)	99.4 (98.3, 99.9)	91.3 (88.5, 93.6)
18-64	Adacel	742-743	97.3 (95.9, 98.3)	72.9 (69.6, 76.1)	100.0 [‡] (99.5, 100.0)	97.8 [§] (96.5, 98.8)	63.1 [‡] (59.5, 66.6)
	Td**	509	95.9 (93.8, 97.4)	70.3 (66.2, 74.3)	99.8 (98.9, 100.0)	98.2 (96.7, 99.2)	66.8 (62.5, 70.9)

^{*} N = number of participants in the per-protocol population with available data.

Booster response is defined as: A four-fold rise in antibody concentration, if the pre-vaccination concentration was equal to or below the cut-off value and a two-fold rise in antibody concentration if the pre-vaccination concentration was above the cut-off value. The cut-off value for tetanus was 2.7 IU/mL.

Seroprotection rates at ≥0.10 IU/mL and booster response rates to Adacel vaccine were non-inferior to Td vaccine (upper limit of the 95% CI on the difference for Td vaccine minus Adacel vaccine <10%).

Seroprotection rates at ≥ 1.0 IU/mL were not prospectively defined as a primary endpoint.

Tetanus and Diphtheria Toxoids Adsorbed for Adult Use manufactured by Sanofi Pasteur Inc., Swiftwater, PA.

Table 4: Pre-vaccination and Post-vaccination Antibody Responses and Booster Response
Rates to Diphtheria Toxoid Following Adacel Vaccine as Compared to Td Vaccine in
Adolescents and Adults 11 Through 64 Years of Age

			Diphtheria Antitoxin (IU/mL)				
			Pre-vaccination		1 Month Post-vaccination		
Age Group (years)	Vaccine	N*	% ≥0.10 (95% CI)	% ≥1.0 (95% CI)	% ≥0.10 (95% CI)	% ≥1.0 (95% CI)	% Booster [†] (95% CI)
11-17	Adacel	527	72.5 (68.5, 76.3)	15.7 (12.7, 19.1)	99.8 [‡] (98.9, 100.0)	98.7 [§] (97.3, 99.5)	95.1 [‡] (92.9, 96.8)
11-17	Td**	515-516	70.7 (66.5, 74.6)	17.3 (14.1, 20.8)	99.8 (98.9, 100.0)	98.4 (97.0, 99.3)	95.0 (92.7, 96.7)
18-64	Adacel	739-741	62.6 (59.0, 66.1)	14.3 (11.9, 17.0)	94.1 [‡] (92.1, 95.7)	78.0 [§] (74.8, 80.9)	87.4 [‡] (84.8, 89.7)
10-04	Td**	506-507	63.3 (59.0, 67.5)	16.0 (12.9, 19.5)	95.1 (92.8, 96.8)	79.9 (76.1, 83.3)	83.4 (79.9, 86.5)

- * N = number of participants in the per-protocol population with available data.
- Booster response is defined as: A four-fold rise in antibody concentration, if the pre-vaccination concentration was equal to or below the cut-off value and a two-fold rise in antibody concentration if the pre-vaccination concentration was above the cut-off value. The cut-off value for diphtheria was 2.56 IU/mL.
- [‡] Seroprotection rates at ≥0.10 IU/mL and booster response rates to Adacel vaccine were non-inferior to Td vaccine (upper limit of the 95% CI on the difference for Td vaccine minus Adacel vaccine <10%).
- § Seroprotection rates at ≥ 1.0 IU/mL were not prospectively defined as a primary endpoint.
- Tetanus and Diphtheria Toxoids Adsorbed for Adult Use manufactured by Sanofi Pasteur Inc., Swiftwater, PA.

387 388

389

Table 5: Ratio of Pertussis Antibody Geometric Mean Concentrations (GMCs)[¥] Observed One Month After a Dose of Adacel Vaccine in Adolescents and Adults 11 Through 64 Years of Age Compared With Those Observed in Infants One Month Following Vaccination at 2, 4 and 6 Months of Age in the Efficacy Trial With DAPTACEL Vaccine

	Adolescents 11-17 Years of Age	Adults 18-64 Years of Age
	Adacel*/DAPTACEL [†]	$\mathbf{Adacel}^{\ddagger}/\mathbf{DAPTACEL}^{\dagger}$
	GMC Ratio	GMC Ratio
	(95% CIs)	(95% CIs)
Anti-PT	3.6	2.1
Allu-F I	$(2.8, 4.5)^{\S}$	$(1.6, 2.7)^{\S}$
Anti-FHA	5.4	4.8
Anu-rha	$(4.5, 6.5)^{\S}$	$(3.9, 5.9)^{\S}$
A 42 DDNI	3.2	3.2
Anti-PRN	$(2.5, 4.1)^{\S}$	$(2.3, 4.4)^{\S}$
A 42 EVINA	5.3	2.5
Anti-FIM	(3.9, 7.1) [§]	$(1.8, 3.5)^{\S}$

[¥] Antibody GMCs, measured in arbitrary ELISA units were calculated separately for infants, adolescents and adults.

^{*} N = 524 to 526, number of adolescents in the per-protocol population with available data for Adacel vaccine.

N = 80, number of infants who received DAPTACEL vaccine with available data post-dose 3 (Sweden Efficacy I).

 $^{^{\}ddagger}$ N = 741, number of adults in the per-protocol population with available data for Adacel vaccine.

GMC following Adacel vaccine was non-inferior to GMC following DAPTACEL vaccine (lower limit of 95% CI on the ratio of GMC for Adacel vaccine divided by DAPTACEL vaccine >0.67).

391

Table 6: Booster Response Rates to the Pertussis Antigens Observed One Month After a Dose of Adacel Vaccine in Adolescents and Adults 11 Through 64 Years of Age

	Adolescents 11-17 Years of Age		Adults 18-64 Years of Age		Pre-defined - Acceptable Rates*
	\mathbf{N}^{\ddagger}	% (95% CI)	\mathbf{N}^{\ddagger}	% (95% CI)	% [†]
Anti-PT	524	92.0 (89.3, 94.2)	739	84.4 (81.6, 87.0)	81.2
Anti-FHA	526	85.6 (82.3, 88.4)	739	82.7 (79.8, 85.3)	77.6
Anti-PRN	525	94.5 (92.2, 96.3)	739	93.8 (91.8, 95.4)	86.4
Anti-FIM	526	94.9 (92.6, 96.6)	739	85.9 (83.2, 88.4)	82.4

^{*} The acceptable response rate for each antigen was defined as the lower limit of the 95% CI for the rate being no more than 10% lower than the response rate observed in previous clinical trials.

The cut-off values were 85 EU/mL for PT, 170 EU/mL for FHA, 115 EU/mL for PRN and 285 EU/mL for FIM

A booster response for each antigen was defined as a four-fold rise in antibody concentration if the pre-vaccination concentration was equal to or below the cut-off value and a two-fold rise in antibody concentration if the pre-vaccination concentration was above the cut-off value. The cut-off values for pertussis antigens were established based on antibody data from both adolescents and adults in previous clinical trials.

 $^{^{\}dagger}$ N = number of participants in the per-protocol population with available data.

392	Study Td519 assessed the comparative immunogenicity of Adacel administered to adolescents
393	(10 to <11 years of age and 11 to <12 years of age) [see Adverse Reactions (6.1).] In this study
394	non-inferiority was demonstrated for booster responses to tetanus and diphtheria toxoids, GMCs
395	to the pertussis antigens (PT, FHA, PRN and FIM) and booster responses to the pertussis antigens
396	PT, FHA and PRN. For FIM, non-inferiority was not demonstrated as the lower bound of the 95%
397	CI of the difference in booster response rates (-5.96%) did not meet the predefined criterion (>-
398	5% when the booster response in the older age group was >95%).
399	14.2 Concomitant Hepatitis B Vaccine Administration
400	The concomitant use of Adacel vaccine and hepatitis B (Hep B) vaccine (Recombivax HB [®] , 10
401	mcg per dose using a two-dose regimen, manufactured by Merck and Co., Inc) was evaluated in a
402	multi-center, open-labeled, randomized, controlled study that enrolled 410 adolescents, 11
403	through 14 years of age inclusive. One group received Adacel and Hep B vaccines concurrently
404	(N=206). The other group $(N=204)$ received Adacel vaccine at the first visit, then 4-6 weeks
405	later received Hep B vaccine. The second dose of Hep B vaccine was given 4-6 weeks after the
406	first dose. Serum samples were obtained prior to and 4-6 weeks after Adacel vaccine
407	administration, as well as 4-6 weeks after the 2 nd dose of Hep B for all participants. No
408	interference was observed in the immune responses to any of the vaccine antigens when Adacel
409	and Hep B vaccines were given concurrently or separately. [See ADVERSE REACTIONS (6.1).]
410	14.3 Concomitant Influenza Vaccine Administration
411	The concomitant use of Adacel vaccine and trivalent inactivated influenza vaccine (TIV,
412	Fluzone®, manufactured by Sanofi Pasteur Inc., Swiftwater, PA) was evaluated in a multi-center,
413	open-labeled, randomized, controlled study conducted in 720 adults, 19-64 years of age inclusive.
414	In one group, participants received Adacel and TIV vaccines concurrently ($N=359$). The other
415	group received TIV at the first visit, then 4-6 weeks later received Adacel vaccine ($N = 361$). Sera
416	were obtained prior to and 4-6 weeks after Adacel vaccine, as well as 4-6 weeks after the TIV.
417	The immune responses were comparable for concurrent and separate administration of Adacel and
418	TIV vaccines for diphtheria (percent of participants with seroprotective concentration \geq 0.10
419	IU/mL and booster responses), tetanus (percent of participants with seroprotective concentration
420	\geq 0.10 IU/mL), pertussis antigens (booster responses and GMCs except lower PRN GMC in the
421	concomitant group, lower bound of the 90% CI was 0.61 and the pre-specified criterion was

422	\geq 0.67) and influenza antigens (percent of participants with hemagglutination-inhibition [HI]
423	antibody titer $\ge 1:40~IU/mL$ and ≥ 4 -fold rise in HI titer). Although tetanus booster response rates
424	were significantly lower in the group receiving the vaccines concurrently versus separately,
425	greater than 98% of participants in both groups achieved seroprotective levels of \geq 0.1 IU/mL.
426	[See ADVERSE REACTIONS (6.1).]
427	

428 429	15	REFERENCES
430	1	Stratton KR, et al, editors. Adverse events associated with childhood vaccines; evidence
431		bearing on causality. Washington: National Academy Press; 1994. p. 67-117.
432	2	Stainer DW, et al. A simple chemically defined medium for the production of phase I
433		Bordetella pertussis. J Gen Microbiol 1970;63:211-20.
434	3	Mueller JH, et al. Variable factors influencing the production of tetanus toxin. J Bacteriol
435		1954;67(3):271-7.
436	4	Stainer DW. Production of diphtheria toxin. In: Manclark CR, editor. Proceedings of an
437		informal consultation on the World Health Organization requirements for diphtheria,
438		tetanus, pertussis and combined vaccines. United States Public Health Service, Bethesda,
439		MD. DHHS 91-1174. 1991. p. 7-11.
440	5	FDA. Department of Health and Human Services (DHHS). Biological products bacterial
441		vaccines and toxoids; implementation of efficacy review; proposed rule. Fed Reg
442		1985;50(240):51002-117.
443	6	Wassilak SGF, et al. Tetanus toxoid. In: Plotkin SA, Orenstein WA, Offit PA, editors.
444		Vaccines. 5th ed. Philadelphia, PA: W.B. Saunders Company; 2008. p. 805-39.
445	7	Vitek CR and Wharton M. Diphtheria toxoid. In: Plotkin SA, Orenstein WA, Offit PA,
446		editors. Vaccines. 5th ed. Philadelphia, PA: W.B. Saunders Company; 2008. p. 139-56.
447	8	Gustafsson L, et al. A controlled trial of a two-component acellular, a five-component
448		acellular and a whole-cell pertussis vaccine. N Engl J Med 1996;334(6):349-55.
449		
450		

451	16 HOW SUPPLIED/STORAGE AND HANDLING
452	Syringe, without needle, 1 dose - NDC No. 49281-400-89 (not made with natural rubber latex); in
453	package of 5 syringes, NDC No. 49281-400-20.
454	Syringe, without needle, 1 dose - NDC No. 49281-400-88; in package of 5 syringes, NDC No.
455	49281-400-15. The tip caps of the prefilled syringes may contain natural rubber latex. No other
456	components are made with natural rubber latex.
457	Vial, 1 dose - NDC No. 49281-400-58; in package of 5 vials; NDC No. 49281-400-05. The vial
458	stopper is not made with natural rubber latex.
459	Vial, 1 dose - NDC No. 49281-400-58; in package of 10 vials; NDC No. 49281-400-10. The vial
460	stopper is not made with natural rubber latex.
461	Adacel vaccine should be stored at 2° to 8°C (35° to 46°F). DO NOT FREEZE. Product which
462	has been exposed to freezing should not be used. Do not use after expiration date shown on the
463	label.
464	17 PATIENT COUNSELING INFORMATION
465	Before administration of Adacel vaccine, health-care providers should inform the patient, parent
466	or guardian of the benefits and risks of the vaccine and the importance of receiving recommended
467	booster dose unless a contraindication to further immunization exists.
468	The health-care provider should inform the patient, parent or guardian about the potential for
469	adverse reactions that have been temporally associated with Adacel vaccine or other vaccines
470	containing similar components. The health-care provider should provide the Vaccine Information
471	Statements (VISs) that are required by the National Childhood Vaccine Injury Act of 1986 to be
472	given with each immunization. The patient, parent or guardian should be instructed to report any
473	serious adverse reactions to their health-care provider.
474	Pregnancy Exposure Registry [See USE IN SPECIFIC POPULATIONS (8.1).]
475	
476	Printed in XXX
477	Manufactured by:
478	Sanofi Pasteur Limited
479	Toronto Ontario Canada

Sanofi	Pasteur
306 - A	\dacel [®]

Full Prescribing Information

481	Distributed by:	
482	Sanofi Pasteur Inc.	
483	Swiftwater PA 18370 USA	
484		
485	Adacel® is a registered trademark of Sanofi, its affiliates and its subsidiaries.	
486		R10-XX17 USA

SANOFI PASTEUR 🔊