

1 **HIGHLIGHTS OF PRESCRIBING INFORMATION**

2 **These highlights do not include all the information needed to use**
3 **FLUMIST® QUADRIVALENT safely and effectively. See full prescribing**
4 **information for FluMist® Quadrivalent.**

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6 **FluMist® Quadrivalent (Influenza Vaccine Live, Intranasal)**
7 **Intranasal Spray**
8 **2015-2016 Formula**
9 **Initial U.S. Approval: 2003**

11 -----INDICATIONS AND USAGE-----

12 FluMist Quadrivalent is a vaccine indicated for active immunization for the
13 prevention of influenza disease caused by influenza A subtype viruses and
14 type B viruses contained in the vaccine. (1, 11)
15 FluMist Quadrivalent is approved for use in persons 2 through 49 years of
16 age. (1)

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

18 For intranasal administration by a healthcare provider. (2)

Age	Dose	Schedule
2 years through 8 years	1 or 2 doses ^a , 0.2 mL ^b each	If 2 doses, administer at least 1 month apart
9 years through 49 years	1 dose, 0.2 mL ^b	-

19 ^a 1 or 2 doses depends on vaccination history as per Advisory Committee on
20 Immunization Practices annual recommendations on prevention and control of
21 influenza with vaccines.

22 ^b Administer as 0.1 mL per nostril.

23 “-” indicates information is not applicable

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

25 Each 0.2 mL dose is a suspension supplied in a single-dose pre-filled
26 intranasal sprayer. (3)

27 -----CONTRAINDICATIONS-----

- 28 • Severe allergic reaction (e.g., anaphylaxis) to any component of FluMist
29 Quadrivalent, including egg protein, or after a previous dose of any
30 influenza vaccine. (4.1, 11)
- 31 • Concomitant aspirin therapy in children and adolescents. (4.2)

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

- 36 • In clinical trials, risks of hospitalization and wheezing were increased in
37 children younger than 2 years of age who received FluMist (trivalent
38 Influenza Vaccine Live, Intranasal). (5.1)
- 39 • Children younger than 5 years of age with recurrent wheezing and persons
40 of any age with asthma may be at increased risk of wheezing following the
41 administration of FluMist Quadrivalent. (5.2)
- 42 • If Guillain-Barré syndrome has occurred within 6 weeks of any prior
43 influenza vaccination, the decision to give FluMist Quadrivalent should be
44 based on careful consideration of the potential benefits and risks. (5.3)
- 45 • FluMist Quadrivalent has not been studied in immunocompromised
46 persons. (5.4)

47 -----ADVERSE REACTIONS-----

48 The most common solicited adverse reactions (≥ 10% in vaccine recipients
49 and at least 5% greater than in placebo recipients) reported after FluMist were
50 runny nose or nasal congestion (ages 2 years through 49 years), fever over
51 100°F (children ages 2 years through 6 years), and sore throat (adults ages 18
52 years through 49 years). Among children and adolescents 2 through 17 years
53 of age who received FluMist Quadrivalent, 32% reported runny nose or nasal
54 congestion and 7% reported fever over 100°F. Among adults 18 through
55 49 years of age who received FluMist Quadrivalent, 44% reported runny nose
56 or nasal congestion and 19% reported sore throat. (6.1)

57 **To report SUSPECTED ADVERSE REACTIONS, contact MedImmune
58 at 1-877-633-4411 or VAERS at 1-800-822-7967 or <http://vaers.hhs.gov>.**

59 -----DRUG INTERACTIONS-----

- 60 • Antiviral drugs that are active against influenza A and/or B may reduce the
61 effectiveness of FluMist Quadrivalent if administered within 48 hours
62 before, or within 2 weeks after, receipt of the vaccine. (7.2)

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

- 64 • Safety and effectiveness of FluMist Quadrivalent have not been established
65 in pregnant women, nursing mothers, geriatric adults, or children less than
66 2 years of age. (8.1, 8.3, 8.4, 8.5)
- 67 • In clinical trials, in children 6 through 23 months of age, FluMist was
68 associated with an increased risk of hospitalization and wheezing. (8.4)

69 **See 17 for PATIENT COUNSELING INFORMATION and FDA-
70 approved patient labeling.**

71 **Revised: 7/2015**

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139 listed.

140 **FULL PRESCRIBING INFORMATION**

141 **1 INDICATIONS AND USAGE**

142 FluMist® Quadrivalent is a vaccine indicated for active immunization for the prevention of influenza
143 disease caused by influenza A subtype viruses and type B viruses contained in the vaccine [see
144 *Description (11)*].

145 FluMist Quadrivalent is approved for use in persons 2 through 49 years of age.

146 **2 DOSAGE AND ADMINISTRATION**

147 **FOR INTRANASAL ADMINISTRATION BY A HEALTHCARE PROVIDER.**

148 **2.1 Dosing Information**

149 Administer FluMist Quadrivalent according to the following schedule:

Age	Dose	Schedule
2 years through 8 years	1 or 2 doses ^a , 0.2 mL ^b each	If 2 doses, administer at least 1 month apart
9 years through 49 years	1 dose, 0.2 mL ^b	-

150 ^a 1 or 2 doses depends on vaccination history as per Advisory Committee on Immunization Practices annual
151 recommendations on prevention and control of influenza with vaccines.

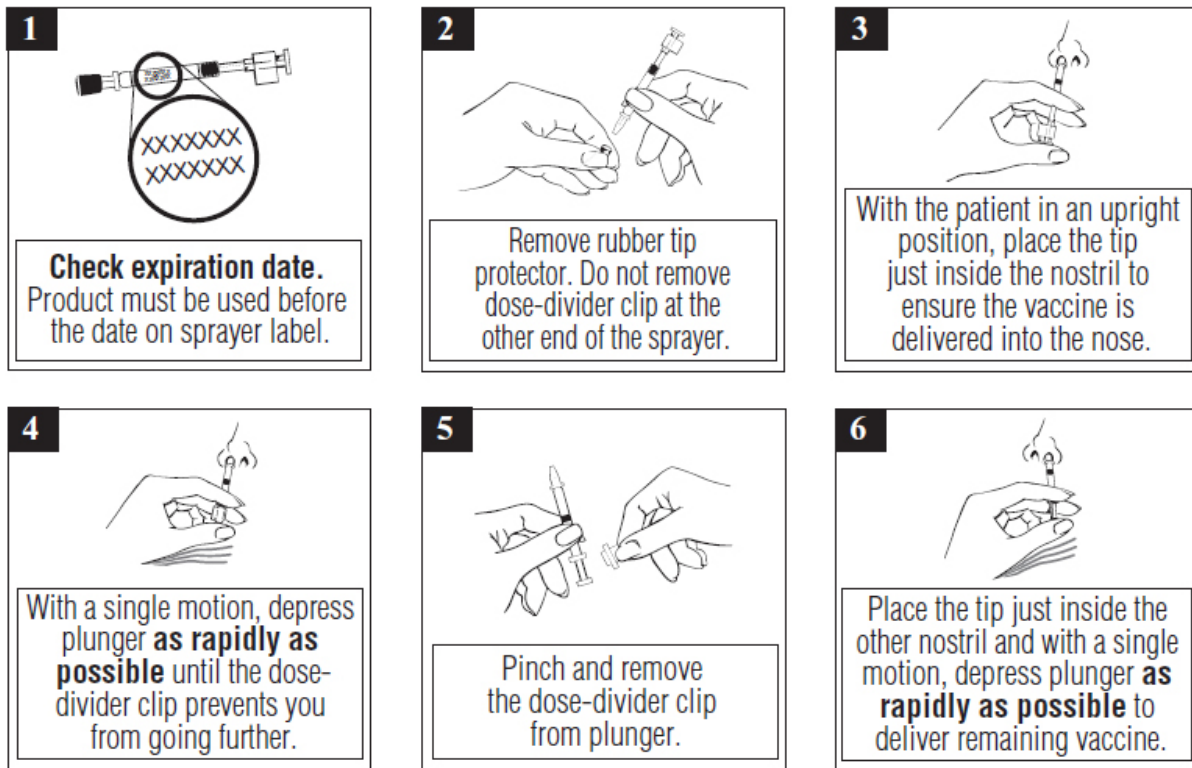
152 ^b Administer as 0.1 mL per nostril.

153 “-” indicates information is not applicable

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155 **2.2 Administration Instructions**

156 Each sprayer contains a single dose (0.2 mL) of FluMist Quadrivalent; administer approximately one half
157 of the contents of the single-dose intranasal sprayer into each nostril (each sprayer contains 0.2 mL of
158 vaccine). Refer to Figure 1 for step-by-step administration instructions. Following administration, dispose
159 of the sprayer according to the standard procedures for medical waste (e.g., sharps container or
160 biohazard container).



 **DO NOT INJECT. DO NOT USE A NEEDLE.**

Note: Active inhalation (i.e., sniffing) is not required by the patient during vaccine administration.

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163 **3 DOSAGE FORMS AND STRENGTHS**

164 Each 0.2 mL dose is a suspension supplied in a single-dose pre-filled intranasal sprayer.

165 **4 CONTRAINDICATIONS**

166 **4.1 Severe Allergic Reactions**

167 Do not administer FluMist Quadrivalent to persons who have had a severe allergic reaction (e.g.,
168 anaphylaxis) to any component of the vaccine [see *Description (11)*] including egg protein, or after a
169 previous dose of any influenza vaccine.

170 **4.2 Concomitant Aspirin Therapy and Reye's Syndrome in Children and Adolescents**

171 Do not administer FluMist Quadrivalent to children and adolescents through 17 years of age who are
172 receiving aspirin therapy or aspirin-containing therapy because of the association of Reye's syndrome
173 with aspirin and wild-type influenza infection [See *Drug Interactions (7.1)*].

174 **5 WARNINGS AND PRECAUTIONS**

175 **5.1 Risks of Hospitalization and Wheezing in Children Younger than 24 Months of Age**

176 In clinical trials, risks of hospitalization and wheezing were increased in children younger than 2 years of
177 age who received FluMist (trivalent Influenza Vaccine Live, Intranasal) [see *Adverse Reactions (6.1)*].
178 This observation with FluMist is relevant to FluMist Quadrivalent because both vaccines are
179 manufactured using the same process and have overlapping compositions [see *Description (11)*].

180 **5.2 Asthma, Recurrent Wheezing, and Active Wheezing**

181 Children younger than 5 years of age with recurrent wheezing and persons of any age with asthma may
182 be at increased risk of wheezing following administration of FluMist Quadrivalent. FluMist Quadrivalent
183 has not been studied in persons with severe asthma or active wheezing.

184 **5.3 Guillain-Barré Syndrome**

185 The 1976 swine influenza vaccine (inactivated) was associated with an elevated risk of Guillain-Barré
186 syndrome (GBS). Evidence for causal relation of GBS with other influenza vaccines is inconclusive; if an
187 excess risk exists, based on data for inactivated influenza vaccines, it is probably slightly more than 1
188 additional case per 1 million persons vaccinated [1]. If GBS has occurred within 6 weeks of any prior
189 influenza vaccination, the decision to give FluMist Quadrivalent should be based on careful consideration
190 of the potential benefits and potential risks.

191 **5.4 Altered Immunocompetence**

192 FluMist Quadrivalent has not been studied in immunocompromised persons. The effectiveness of FluMist
193 has not been studied in immunocompromised persons. Data on safety and shedding of vaccine virus after
194 administration of FluMist in immunocompromised persons are limited to 173 persons with HIV infection
195 and 10 mild to moderately immunocompromised children and adolescents with cancer [see *Clinical*
196 *Pharmacology (12.2)*].

197 **5.5 Medical Conditions Predisposing to Influenza Complications**

198 The safety of FluMist Quadrivalent in individuals with underlying medical conditions that may predispose
199 them to complications following wild-type influenza infection has not been established.

200 **5.6 Management of Acute Allergic Reactions**

201 Appropriate medical treatment and supervision must be available to manage possible anaphylactic
202 reactions following administration of the vaccine [see *Contraindications (4.1)*].

203 **5.7 Limitations of Vaccine Effectiveness**

204 FluMist Quadrivalent may not protect all individuals receiving the vaccine.

6 ADVERSE REACTIONS

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.

This safety experience with FluMist is relevant to FluMist Quadrivalent because both vaccines are manufactured using the same process and have overlapping compositions [see *Description (11)*]. A total of 9537 children and adolescents 1 through 17 years of age and 3041 adults 18 through 64 years of age received FluMist in randomized, placebo-controlled Studies D153-P501, AV006, D153-P526, AV019, and AV009 [3 used Allantoic Fluid containing Sucrose-Phosphate-Glutamate (AF-SPG) placebo, and 2 used saline placebo] described below. In addition, 4179 children 6 through 59 months of age received FluMist in Study MI-CP111, a randomized, active-controlled trial. Among pediatric FluMist recipients 6 months through 17 years of age, 50% were female; in the study of adults, 55% were female. In MI-CP111, AV006, D153-P526, AV019, and AV009, subjects were White (71%), Hispanic (11%), Asian (7%), Black (6%), and Other (5%), while in D153-P501, 99% of subjects were Asian.

A total of 1382 children and adolescents 2 through 17 years of age and 1198 adults 18 through 49 years of age received FluMist Quadrivalent in randomized, active-controlled Studies MI-CP208 and MI-CP185. Among pediatric FluMist Quadrivalent recipients 2 through 17 years of age, 51% were female; in the study of adults, 55% were female. In Studies MI-CP208 and MI-CP185, subjects were White (73%), Asian (1%), Black or African-American (19%), and Other (7%); overall, 22% were Hispanic or Latino.

FluMist in Children and Adolescents

The safety of FluMist was evaluated in an AF-SPG placebo-controlled study (AV019) conducted in a Health Maintenance Organization (HMO) in children 1 through 17 years of age (FluMist = 6473, placebo = 3216). An increase in asthma events, captured by review of diagnostic codes, was observed in children younger than 5 years of age who received FluMist compared to those who received placebo (Relative Risk 3.53, 90% CI: 1.1, 15.7).

In Study MI-CP111, children 6 through 59 months of age were randomized to receive FluMist or inactivated Influenza Virus Vaccine manufactured by Sanofi Pasteur Inc. Wheezing requiring bronchodilator therapy or accompanied by respiratory distress or hypoxia was prospectively monitored from randomization through 42 days post last vaccination. Hospitalization due to all causes was prospectively monitored from randomization through 180 days post last vaccination. Increases in wheezing and hospitalization (for any cause) were observed in children 6 months through 23 months of age who received FluMist compared to those who received inactivated Influenza Virus Vaccine, as shown in Table 1.

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Table 1: Percentages of Children with Hospitalizations and Wheezing from Study MI-CP111^a

Adverse Reaction	Age Group	FluMist (n/N)	Active Control^b (n/N)
Hospitalizations ^c	6-23 months	4.2% (84/1992)	3.2% (63/1975)
	24-59 months	2.1% (46/2187)	2.5% (56/2198)
Wheezing ^d	6-23 months	5.9% (117/1992)	3.8% (75/1975)
	24-59 months	2.1% (47/2187)	2.5% (56/2198)

240 ^a NCT00128167; see www.clinicaltrials.gov241 ^b Inactivated Influenza Virus Vaccine manufactured by Sanofi Pasteur Inc., administered intramuscularly.242 ^c Hospitalization due to any cause from randomization through 180 days post last vaccination.243 ^d Wheezing requiring bronchodilator therapy or accompanied by respiratory distress or hypoxia evaluated from
244 randomization through 42 days post last vaccination.

245 Most hospitalizations observed were due to gastrointestinal and respiratory tract infections and occurred
246 more than 6 weeks post vaccination. In post-hoc analysis, rates of hospitalization in children 6 through
247 11 months of age were 6.1% (42/684) in FluMist recipients and 2.6% (18/683) in inactivated Influenza
248 Virus Vaccine recipients.

249 Table 2 shows pooled solicited adverse reactions occurring in at least 1% of FluMist recipients and at a
250 higher rate ($\geq 1\%$ rate difference after rounding) compared to placebo post Dose 1 for Studies D153-P501
251 and AV006, and solicited adverse reactions post Dose 1 for Study MI-CP111. Solicited adverse reactions
252 were those about which parents/guardians were specifically queried after receipt of FluMist, placebo, or
253 control vaccine. In these studies, solicited reactions were documented for 10 days post vaccination.
254 Solicited reactions following the second dose of FluMist were similar to those following the first dose and
255 were generally observed at a lower frequency.

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Table 2: Summary of Solicited Adverse Reactions Observed Within 10 Days after Dose 1 for FluMist and Either Placebo or Active Control Recipients in Children 2 through 6 Years of Age

Event	Studies D153-P501 ^a & AV006		Study MI-CP111 ^b	
	FluMist N = 876-1759 ^e	Placebo ^c N = 424-1034 ^e	FluMist N = 2170 ^e	Active Control ^d N = 2165 ^e
	%	%	%	%
Runny Nose/ Nasal Congestion	58	50	51	42
Decreased Appetite	21	17	13	12
Irritability	21	19	12	11
Decreased Activity (Lethargy)	14	11	7	6
Sore Throat	11	9	5	6
Headache	9	7	3	3
Muscle Aches	6	3	2	2
Chills	4	3	2	2
Fever				
> 100°F Oral	16	11	13	11
> 100 - ≤ 101°F Oral	9	6	6	4
> 101 - ≤ 102°F Oral	4	3	4	3

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^a NCT00192244; see www.clinicaltrials.gov
^b NCT00128167; see www.clinicaltrials.gov
^c Study D153-P501 used saline placebo; Study AV006 used AF-SPG placebo.
^d Inactivated Influenza Virus Vaccine manufactured by Sanofi Pasteur Inc., administered intramuscularly.
^e Number of evaluable subjects (those who returned diary cards) for each reaction. Range reflects differences in data collection between the 2 pooled studies.

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In clinical studies D153-P501 and AV006, unsolicited adverse reactions in children occurring in at least 1% of FluMist recipients and at a higher rate (≥ 1% rate difference after rounding) compared to placebo were abdominal pain (2% FluMist vs. 0% placebo) and otitis media (3% FluMist vs. 1% placebo). An additional adverse reaction identified in the active-controlled trial MI-CP111 occurring in at least 1% of FluMist recipients and at a higher rate (≥ 1% rate difference after rounding) compared to active control was sneezing (2% FluMist vs. 1% active control).

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In a separate saline placebo-controlled trial (D153-P526) in a subset of older children and adolescents 9 through 17 years of age who received one dose of FluMist, the solicited adverse reactions as well as unsolicited adverse reactions reported were generally consistent with observations from the trials in Table 2. Abdominal pain was reported in 12% of FluMist recipients compared to 4% of placebo recipients and decreased activity was reported in 6% of FluMist recipients compared to 0% of placebo recipients.

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In Study AV018, in which FluMist was concomitantly administered with Measles, Mumps, and Rubella Virus Vaccine Live (MMR, manufactured by Merck & Co., Inc.) and Varicella Virus Vaccine Live (manufactured by Merck & Co., Inc.) to children 12 through 15 months of age, adverse reactions were similar to those seen in other clinical trials of FluMist.

FluMist Quadrivalent in Children and Adolescents

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In the randomized, active-controlled Study MI-CP208 that compared FluMist Quadrivalent and FluMist in children and adolescents 2 through 17 years of age, the rates of solicited adverse reactions reported

282 were similar between subjects who received FluMist Quadrivalent and FluMist. Table 3 includes solicited
 283 adverse reactions post Dose 1 from Study MI-CP208 that either occurred at a higher rate ($\geq 1\%$ rate
 284 difference after rounding) in FluMist Quadrivalent recipients compared to FluMist recipients or were
 285 identified in previous FluMist clinical studies (see *Table 2*). In this study, solicited adverse reactions were
 286 documented for 14 days post vaccination. Solicited adverse reactions post Dose 2 were observed at a
 287 lower frequency compared to those post Dose 1 for FluMist Quadrivalent and were similar between
 288 subjects who received FluMist Quadrivalent and FluMist.

289 **Table 3: Summary of Solicited Adverse Reactions^a Observed Within 14 Days after Dose 1 for**
 290 **FluMist Quadrivalent and FluMist Recipients in Study MI-CP208^b in Children and Adolescents 2**
 291 **through 17 Years of Age**

	FluMist Quadrivalent N = 1341-1377 ^d	FluMist ^c N = 901-920 ^d
Event	%	%
Runny Nose/Nasal Congestion	32	32
Headache	13	12
Decreased Activity (Lethargy)	10	10
Sore Throat	9	10
Decreased Appetite	6	7
Muscle Aches	4	5
Fever		
> 100°F by any route	7	5
> 100 - \leq 101°F by any route	3	2
> 101 - \leq 102°F by any route	2	2

292 ^a Solicited adverse reactions that occurred at a higher rate ($\geq 1\%$ rate difference after rounding) in
 293 FluMist Quadrivalent recipients compared to FluMist recipients or were identified in previous FluMist
 294 trials (see *Table 2*).

295 ^b NCT01091246; see www.clinicaltrials.gov

296 ^c Represents pooled data from the two FluMist study arms. [see *Clinical Studies (14.2)*]

297 ^d Number of evaluable subjects for each event.

298 In Study MI-CP208, no unsolicited adverse reactions occurred at a higher rate (1% or greater) in FluMist
 299 Quadrivalent recipients compared to FluMist recipients.

300 **FluMist in Adults**

301 In adults 18 through 49 years of age in Study AV009, solicited adverse reactions occurring in at least
 302 1% of FluMist recipients and at a higher rate ($\geq 1\%$ rate difference after rounding) compared to AF-SPG
 303 placebo include runny nose (44% FluMist vs. 27% placebo), headache (40% FluMist vs. 38% placebo),
 304 sore throat (28% FluMist vs. 17% placebo), tiredness/weakness (26% FluMist vs. 22% placebo), muscle
 305 aches (17% FluMist vs. 15% placebo), cough (14% FluMist vs. 11% placebo), and chills (9% FluMist vs.
 306 6% placebo).

307 In Study AV009, unsolicited adverse reactions occurring in at least 1% of FluMist recipients and at a
 308 higher rate ($\geq 1\%$ rate difference after rounding) compared to placebo were nasal congestion (9% FluMist
 309 vs. 2% placebo) and sinusitis (4% FluMist vs. 2% placebo).

310 **FluMist Quadrivalent in Adults**

311 In the randomized, active-controlled Study MI-CP185 that compared FluMist Quadrivalent and FluMist in
312 adults 18 through 49 years of age, the rates of solicited adverse reactions reported were generally similar
313 between subjects who received FluMist Quadrivalent and FluMist. Table 4 presents solicited adverse
314 reactions that either occurred at a higher rate ($\geq 1\%$ rate difference after rounding) in FluMist Quadrivalent
315 recipients compared to FluMist recipients or were identified in Study AV009.

316 **Table 4: Summary of Solicited Adverse Reactions^a Observed Within 14 Days after Dose 1 for**
317 **FluMist Quadrivalent and FluMist Recipients in Study MI-CP185^b in Adults 18 through 49 Years of**
318 **Age**

	FluMist Quadrivalent N = 1197^d	FluMist^c N = 597^d
Event	%	%
Runny Nose/Nasal Congestion	44	40
Headache	28	27
Sore Throat	19	20
Decreased Activity (Lethargy)	18	18
Cough	14	13
Muscle Aches	10	10
Decreased Appetite	6	5

319 ^a Solicited adverse reactions that occurred at a higher rate ($\geq 1\%$ rate difference after
320 rounding) in FluMist Quadrivalent recipients compared to FluMist recipients or were
321 identified in Study AV009.

322 ^b NCT00860067; see www.clinicaltrials.gov

323 ^c Represents pooled data from the two FluMist study arms. [see *Clinical Studies (14.4)*]

324 ^d Number of evaluable subjects for each event.

325 In Study MI-CP185, no unsolicited adverse reactions occurred at a higher rate (1% or greater) in FluMist
326 Quadrivalent recipients compared to FluMist recipients.

327 **6.2 Postmarketing Experience**

328 The following events have been spontaneously reported during post approval use of FluMist. Because
329 these events are reported voluntarily from a population of uncertain size, it is not always possible to
330 reliably estimate their frequency or establish a causal relationship to vaccine exposure.

331 Cardiac disorders: Pericarditis

332 Congenital, familial, and genetic disorders: Exacerbation of symptoms of mitochondrial
333 encephalomyopathy (Leigh syndrome)

334 Gastrointestinal disorders: Nausea, vomiting, diarrhea

335 Immune system disorders: Hypersensitivity reactions (including anaphylactic reaction, facial edema, and
336 urticaria)

337 Nervous system disorders: Guillain-Barré syndrome, Bell's Palsy, meningitis, eosinophilic meningitis,
338 vaccine-associated encephalitis

339 Respiratory, thoracic, and mediastinal disorders: Epistaxis

340 Skin and subcutaneous tissue disorders: Rash

341 **7 DRUG INTERACTIONS**

342 **7.1 Aspirin Therapy**

343 Do not administer FluMist Quadrivalent to children and adolescents through 17 years of age who are
344 receiving aspirin therapy or aspirin-containing therapy because of the association of Reye's syndrome
345 with aspirin and wild-type influenza [see *Contraindications (4.2)*]. Avoid aspirin-containing therapy in
346 these age groups during the first 4 weeks after vaccination with FluMist Quadrivalent unless clearly
347 needed.

348 **7.2 Antiviral Agents Against Influenza A and/or B**

349 Antiviral drugs that are active against influenza A and/or B viruses may reduce the effectiveness of
350 FluMist Quadrivalent if administered within 48 hours before, or within 2 weeks after vaccination. The
351 concurrent use of FluMist Quadrivalent with antiviral agents that are active against influenza A and/or
352 B viruses has not been evaluated. If antiviral agents and FluMist Quadrivalent are administered
353 concomitantly, revaccination should be considered when appropriate.

354 **7.3 Concomitant Administration with Inactivated Vaccines**

355 The safety and immunogenicity of FluMist Quadrivalent when administered concomitantly with inactivated
356 vaccines have not been determined. Studies of FluMist and FluMist Quadrivalent excluded subjects who
357 received any inactivated or subunit vaccine within two weeks of enrollment.

358 **7.4 Concomitant Administration with Other Live Vaccines**

359 Concomitant administration of FluMist Quadrivalent with Measles, Mumps, and Rubella Virus Vaccine
360 Live (MMR, manufactured by Merck & Co., Inc.) or the Varicella Virus Vaccine Live (manufactured by
361 Merck & Co., Inc.) has not been studied. Concomitant administration of FluMist with MMR and the
362 varicella vaccine was studied in children 12 through 15 months of age [see *Clinical Studies (14.5)*].
363 Concomitant administration of FluMist with the MMR and the varicella vaccine in children older than
364 15 months of age has not been studied.

365 **7.5 Intranasal Products**

366 There are no data regarding co-administration of FluMist Quadrivalent with other intranasal preparations.

367 **8 USE IN SPECIFIC POPULATIONS**

368 **8.1 Pregnancy**

369 **Pregnancy Category B**

370 A developmental and reproductive toxicity study has been performed in female rats administered FluMist
371 Quadrivalent either three times (during the period of organogenesis) or six times (prior to gestation and

372 during the period of organogenesis), 200 microliter/rat/occasion (approximately 150 human dose
373 equivalents), by intranasal instillation and has revealed no evidence of impaired fertility or harm to the
374 fetus due to FluMist Quadrivalent. There are however, no adequate and well controlled studies in
375 pregnant women. Because animal studies are not always predictive of human response FluMist
376 Quadrivalent should be administered during pregnancy only if clearly needed.

377 **8.3 Nursing Mothers**

378 It is not known whether FluMist Quadrivalent is excreted in human milk. Because some viruses are
379 excreted in human milk, caution should be exercised when FluMist Quadrivalent is administered to a
380 nursing woman.

381 **8.4 Pediatric Use**

382 Safety and effectiveness of FluMist Quadrivalent in children 24 months of age and older is based on data
383 from FluMist clinical studies and a comparison of post-vaccination antibody titers between persons who
384 received FluMist Quadrivalent and those who received FluMist [see *Clinical Studies (14.1, 14.2)*]. FluMist
385 Quadrivalent is not approved for use in children younger than 24 months of age because use of FluMist in
386 children 6 through 23 months has been associated with increased risks of hospitalization and wheezing in
387 clinical trials [see *Warnings and Precautions (5.1)* and *Adverse Reactions (6.1)*].

388 **8.5 Geriatric Use**

389 FluMist Quadrivalent is not approved for use in persons 65 years of age and older because in a clinical
390 study (AV009), effectiveness of FluMist to prevent febrile illness was not demonstrated in adults
391 50 through 64 years of age [see *Clinical Studies (14.3)*]. In this study, solicited events among individuals
392 50 through 64 years of age were similar in type and frequency to those reported in younger adults. In a
393 clinical study of FluMist in persons 65 years of age and older, subjects with underlying high-risk medical
394 conditions (N = 200) were studied for safety. Compared to controls, FluMist recipients had a higher rate of
395 sore throat.

396 **11 DESCRIPTION**

397 FluMist Quadrivalent (Influenza Vaccine Live, Intranasal) is a live quadrivalent vaccine for administration
398 by intranasal spray. FluMist Quadrivalent contains four vaccine virus strains: an A/H1N1 strain, an
399 A/H3N2 strain and two B strains. FluMist Quadrivalent contains B strains from both the
400 B/Yamagata/16/88 and the B/Victoria/2/87 lineages. FluMist Quadrivalent is manufactured according to
401 the same process as FluMist.

402 The influenza virus strains in FluMist Quadrivalent are (a) *cold-adapted (ca)* (i.e., they replicate efficiently
403 at 25°C, a temperature that is restrictive for replication of many wild-type influenza viruses);
404 (b) *temperature-sensitive (ts)* (i.e., they are restricted in replication at 37°C (Type B strains) or 39°C (Type
405 A strains), temperatures at which many wild-type influenza viruses grow efficiently); and (c) *attenuated*

406 (*att*) (i.e., they do not produce classic influenza-like illness in the ferret model of human influenza
407 infection).

408 No evidence of reversion has been observed in the recovered vaccine strains that have been tested
409 (135 of possible 250 recovered isolates) using FluMist [see *Clinical Pharmacology (12.2)*]. For each of the
410 four reassortant strains in FluMist Quadrivalent, the six internal gene segments responsible for *ca*, *ts*, and
411 *att* phenotypes are derived from a master donor virus (MDV), and the two segments that encode the two
412 surface glycoproteins, hemagglutinin (HA) and neuraminidase (NA), are derived from the corresponding
413 antigenically relevant wild-type influenza viruses. Thus, the four viruses contained in FluMist Quadrivalent
414 maintain the replication characteristics and phenotypic properties of the MDV and express the HA and NA
415 of wild-type viruses. For the Type A MDV, at least five genetic loci in three different internal gene
416 segments contribute to the *ts* and *att* phenotypes. For the Type B MDV, at least three genetic loci in two
417 different internal gene segments contribute to both the *ts* and *att* properties; five genetic loci in three gene
418 segments control the *ca* property.

419 Each of the reassortant strains in FluMist Quadrivalent express the HA and NA of wild- type viruses that
420 are related to strains expected to circulate during the 2015-2016 influenza season. Three of the viruses
421 (A/H1N1, A/H3N2 and one B strain) have been recommended by the United States Public Health Service
422 (USPHS) for inclusion in the annual trivalent and quadrivalent influenza vaccine formulations. An
423 additional B strain has been recommended by the USPHS for inclusion in the quadrivalent influenza
424 vaccine formulation.

425 Specific pathogen-free (SPF) eggs are inoculated with each of the reassortant strains and incubated to
426 allow vaccine virus replication. The allantoic fluid of these eggs is harvested, pooled, and then clarified by
427 filtration. The virus is concentrated by ultracentrifugation and diluted with stabilizing buffer to obtain the
428 final sucrose and potassium phosphate concentrations. The viral harvests are then sterile filtered to
429 produce the monovalent bulks. Each lot is tested for *ca*, *ts*, and *att* phenotypes and is also tested
430 extensively by *in vitro* and *in vivo* methods to detect adventitious agents. Monovalent bulks from the four
431 strains are subsequently blended and diluted as required to attain the desired potency with stabilizing
432 buffers to produce the quadrivalent bulk vaccine. The bulk vaccine is then filled directly into individual
433 sprayers for nasal administration.

434 Each pre-filled refrigerated FluMist Quadrivalent sprayer contains a single 0.2 mL dose. Each 0.2 mL
435 dose contains $10^{6.5-7.5}$ FFU (fluorescent focus units) of live attenuated influenza virus reassortants of each
436 of the four strains: A/Bolivia/559/2013 (H1N1) (an A/California/7/2009 (H1N1)pdm09-like virus),
437 A/Switzerland/9715293/2013 (H3N2), B/Phuket/3073/2013 (B/Yamagata/16/88 lineage), and
438 B/Brisbane/60/2008 (B/Victoria/2/87 lineage). Each 0.2 mL dose also contains 0.188 mg/dose
439 monosodium glutamate, 2.00 mg/dose hydrolyzed porcine gelatin, 2.42 mg/dose arginine, 13.68 mg/dose
440 sucrose, 2.26 mg/dose dibasic potassium phosphate, and 0.96 mg/dose monobasic potassium
441 phosphate. Each dose contains residual amounts of ovalbumin (< 0.24 mcg/dose), and may also contain

442 residual amounts of gentamicin sulfate (< 0.015 mcg/mL), and ethylenediaminetetraacetic acid (EDTA)
443 (< 0.37 mcg/dose). FluMist Quadrivalent contains no preservatives.

444 The tip attached to the sprayer is equipped with a nozzle that produces a fine mist that is primarily
445 deposited in the nose and nasopharynx. FluMist Quadrivalent is a colorless to pale yellow suspension
446 and is clear to slightly cloudy.

447 **12 CLINICAL PHARMACOLOGY**

448 **12.1 Mechanism of Action**

449 Immune mechanisms conferring protection against influenza following receipt of FluMist Quadrivalent
450 vaccine are not fully understood; serum antibodies, mucosal antibodies, and influenza-specific T cells
451 may play a role.

452 FluMist and FluMist Quadrivalent contain live attenuated influenza viruses that must infect and replicate in
453 cells lining the nasopharynx of the recipient to induce immunity. Vaccine viruses capable of infection and
454 replication can be cultured from nasal secretions obtained from vaccine recipients (shedding) [see
455 *Pharmacodynamics (12.2)*].

456 **12.2 Pharmacodynamics**

457 **Shedding Studies**

458 Shedding of vaccine viruses within 28 days of vaccination with FluMist was evaluated in (1) multi-center
459 study MI-CP129 which enrolled healthy individuals 6 through 59 months of age (N = 200); and (2) multi-
460 center study FM026 which enrolled healthy individuals 5 through 49 years of age (N = 344). In each
461 study, nasal secretions were obtained daily for the first 7 days and every other day through either Day 25
462 and on Day 28 or through Day 28. In study MI-CP129, individuals with a positive shedding sample at
463 Day 25 or Day 28 were to have additional shedding samples collected every 7 days until culture negative
464 on 2 consecutive samples. Results of these studies are presented in Table 5.

465
466

Table 5: Characterization of Shedding with FluMist in Specified Age Groups by Frequency, Amount, and Duration (Study MI-CP129^a and Study FM026^b)

Age	Number of Subjects	% Shedding ^c	Peak Titer (TCID ₅₀ /mL) ^d	% Shedding After Day 11	Day of Last Positive Culture
6-23 months ^e	99	89	< 5 log ₁₀	7.0	Day 23 ^f
24-59 months	100	69	< 5 log ₁₀	1.0	Day 25 ^g
5-8 years	102	50	< 5 log ₁₀	2.9	Day 23 ^h
9-17 years	126	29	< 4 log ₁₀	1.6	Day 28 ^h
18-49 years	115	20	< 3 log ₁₀	0.9	Day 17 ^h

467 ^a NCT00344305; see www.clinicaltrials.gov
 468 ^b NCT00192140; see www.clinicaltrials.gov
 469 ^c Proportion of subjects with detectable virus at any time point during the 28 days.
 470 ^d Peak titer at any time point during the 28 days among samples positive for a single vaccine virus.
 471 ^e FluMist and FluMist Quadrivalent are not approved for use in children younger than 24 months of age [see
 472 *Adverse Reactions (6.1)*].
 473 ^f A single subject who shed previously on Days 1-3; TCID₅₀/mL was less than 1.5 log₁₀ on Day 23.
 474 ^g A single subject who did not shed previously; TCID₅₀/mL was less than 1.5 log₁₀.
 475 ^h A single subject who did not shed previously; TCID₅₀/mL was less than 1.0 log₁₀.

476 The highest proportion of subjects in each group shed one or more vaccine strains on Days 2-3
 477 post vaccination. After Day 11 among individuals 2 through 49 years of age (n = 443), virus titers did not
 478 exceed 1.5 log₁₀ TCID₅₀/mL.

479 **Studies in Immunocompromised Individuals**

480 Safety and shedding of vaccine virus following FluMist administration were evaluated in 28 HIV-infected
 481 adults [median CD4 cell count of 541 cells/mm³] and 27 HIV-negative adults 18 through 58 years of age.
 482 No serious adverse events were reported during the one-month follow-up period. Vaccine strain (type B)
 483 virus was detected in 1 of 28 HIV-infected subjects on Day 5 only, and in none of the HIV-negative
 484 FluMist recipients.

485 Safety and shedding of vaccine virus following FluMist administration were also evaluated in children in a
 486 randomized (1:1), cross-over, double-blind, AF-SPG placebo-controlled trial in 24 HIV-infected children
 487 [median CD4 cell count of 1013 cells/mm³] and 25 HIV-negative children 1 through 7 years of age, and in
 488 a randomized (1:1), open-label, inactivated influenza vaccine-controlled trial in 243 HIV-infected children
 489 and adolescents 5 through 17 years of age receiving stable anti-retroviral therapy. Frequency and
 490 duration of vaccine virus shedding in HIV-infected individuals were comparable to that seen in healthy
 491 individuals. No adverse effects on HIV viral load or CD4 counts were identified following FluMist
 492 administration. In the 5 through 17 year old age group, one inactivated influenza vaccine recipient and
 493 one FluMist recipient experienced pneumonia within 28 days of vaccination (days 17 and 13,
 494 respectively). The effectiveness of FluMist and FluMist Quadrivalent in preventing influenza illness in HIV-
 495 infected individuals has not been evaluated.

496 Twenty mild to moderately immunocompromised children and adolescents 5 through 17 years of age
 497 (receiving chemotherapy and/or radiation therapy or who had received chemotherapy in the 12 weeks

498 prior to enrollment) were randomized 1:1 to receive FluMist or AF-SPG placebo. Frequency and duration
499 of vaccine virus shedding in these immunocompromised children and adolescents were comparable to
500 that seen in healthy children and adolescents. The effectiveness of FluMist and FluMist Quadrivalent in
501 preventing influenza illness in immunocompromised individuals has not been evaluated.

502 **Transmission Study**

503 A prospective, randomized, double-blind, placebo-controlled trial was performed in a daycare setting in
504 children younger than 3 years of age to assess the transmission of vaccine viruses from a vaccinated
505 individual to a non-vaccinated individual. A total of 197 children 8 through 36 months of age were
506 randomized to receive one dose of FluMist (N = 98) or AF-SPG placebo (N = 99). Virus shedding was
507 evaluated for 21 days by culture of nasal swab specimens. Wild-type A (A/H3N2) influenza virus was
508 documented to have circulated in the community and in the study population during the trial, whereas
509 Type A (A/H1N1) and Type B strains did not.

510 At least one vaccine strain was isolated from 80% of FluMist recipients; strains were recovered from 1-
511 21 days post vaccination (mean duration of 7.6 days \pm 3.4 days). The cold-adapted (*ca*) and temperature-
512 sensitive (*ts*) phenotypes were preserved in 135 tested of 250 strains isolated at the local laboratory. Ten
513 influenza isolates (9 influenza A, 1 influenza B) were cultured from a total of seven placebo subjects. One
514 placebo subject had mild symptomatic Type B virus infection confirmed as a transmitted vaccine virus by
515 a FluMist recipient in the same playgroup. This Type B isolate retained the *ca*, *ts*, and *att* phenotypes of
516 the vaccine strain and had the same genetic sequence when compared to a Type B virus cultured from a
517 vaccine recipient within the same playgroup. Four of the influenza Type A isolates were confirmed as
518 wild-type A/Panama (H3N2). The remaining isolates could not be further characterized.

519 Assuming a single transmission event (isolation of the Type B vaccine strain), the probability of a young
520 child acquiring vaccine virus following close contact with a single FluMist vaccinee in this daycare setting
521 was 0.58% (95% CI: 0, 1.7) based on the Reed-Frost model. With documented transmission of one
522 Type B in one placebo subject and possible transmission of Type A viruses in four placebo subjects, the
523 probability of acquiring a transmitted vaccine virus was estimated to be 2.4% (95% CI: 0.13, 4.6) using
524 the Reed-Frost model.

525 **12.3 Pharmacokinetics**

526 **Biodistribution**

527 A biodistribution study of intranasally administered radiolabeled placebo was conducted in 7 healthy adult
528 volunteers. The mean percentages of the delivered doses detected were as follows: nasal cavity 89.7%,
529 stomach 2.6%, brain 2.4%, and lung 0.4%. The clinical significance of these findings is unknown.

530 **13 NONCLINICAL TOXICOLOGY**

531 **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

532 FluMist Quadrivalent has not been evaluated for its carcinogenic or mutagenic potential or its potential to
533 impair fertility.

534 **14 CLINICAL STUDIES**

535 The effectiveness of FluMist Quadrivalent is based on data demonstrating the clinical efficacy of FluMist
536 in children and the effectiveness of FluMist in adults, and a comparison of post vaccination geometric
537 mean titers (GMTs) of hemagglutination inhibition (HI) antibodies between individuals receiving FluMist
538 and FluMist Quadrivalent. The clinical experience with FluMist is relevant to FluMist Quadrivalent
539 because both vaccines are manufactured using the same process and have overlapping compositions
540 [see *Description (11)*].

541 **14.1 Efficacy Studies of FluMist in Children and Adolescents**

542 A multinational, randomized, double-blind, active-controlled trial (MI-CP111) was performed to assess the
543 efficacy of FluMist compared to an intramuscularly administered, inactivated Influenza Virus Vaccine
544 manufactured by Sanofi Pasteur Inc. (active control) in children 6 months to less than 5 years of age
545 during the 2004-2005 influenza season. A total number of 3916 children without severe asthma, without
546 use of bronchodilator or steroids, and without wheezing within the prior 6 weeks were randomized to
547 FluMist and 3936 were randomized to active control. Children who previously received any influenza
548 vaccine received a single dose of study vaccine, while those who never previously received an influenza
549 vaccination (or had an unknown history of influenza vaccination) received two doses. Participants were
550 then followed through the influenza season to identify illness caused by influenza virus. As the primary
551 endpoint, culture-confirmed modified CDC-ILI (CDC-defined influenza-like illness) was defined as a
552 positive culture for a wild-type influenza virus associated within ± 7 days of modified CDC-ILI. Modified
553 CDC-ILI was defined as fever (temperature $\geq 100^{\circ}\text{F}$ oral or equivalent) with cough, sore throat, or runny
554 nose/nasal congestion on the same or consecutive days.

555 In the primary efficacy analysis, FluMist demonstrated a 44.5% (95% CI: 22.4, 60.6) reduction in influenza
556 rate compared to active control as measured by culture-confirmed modified CDC-ILI caused by wild-type
557 strains antigenically similar to those contained in the vaccine. See Table 6 for a description of the results
558 by strain and antigenic similarity.

Table 6: Comparative Efficacy Against Culture-Confirmed Modified CDC-ILI^a Caused by Wild-Type Strains (Study MI-CP111)^{b,c}

	FluMist			Active Control ^d			% Reduction in Rate for FluMist ^e	95% CI
	N	# of Cases	Rate (cases/N)	N	# of Cases	Rate (cases/N)		
Matched Strains								
All strains	3916	53	1.4%	3936	93	2.4%	44.5%	22.4, 60.6
A/H1N1	3916	3	0.1%	3936	27	0.7%	89.2%	67.7, 97.4
A/H3N2	3916	0	0.0%	3936	0	0.0%	--	--
B	3916	50	1.3%	3936	67	1.7%	27.3%	-4.8, 49.9
Mismatched Strains								
All strains	3916	102	2.6%	3936	245	6.2%	58.2%	47.4, 67.0
A/H1N1	3916	0	0.0%	3936	0	0.0%	--	--
A/H3N2	3916	37	0.9%	3936	178	4.5%	79.2%	70.6, 85.7
B	3916	66	1.7%	3936	71	1.8%	6.3%	-31.6, 33.3
Regardless of Match								
All strains	3916	153	3.9%	3936	338	8.6%	54.9%	45.4, 62.9
A/H1N1	3916	3	0.1%	3936	27	0.7%	89.2%	67.7, 97.4
A/H3N2	3916	37	0.9%	3936	178	4.5%	79.2%	70.6, 85.7
B	3916	115	2.9%	3936	136	3.5%	16.1%	-7.7, 34.7

ATP Population.

560 ^a Modified CDC-ILI was defined as fever (temperature $\geq 100^{\circ}\text{F}$ oral or equivalent) plus cough, sore throat, or runny nose/nasal congestion on the same or
561 consecutive days.

562 ^b In children 6 months through 5 years of age

563 ^c NCT00128167; see www.clinicaltrials.gov

564 ^d Inactivated Influenza Virus Vaccine manufactured by Sanofi Pasteur Inc., administered intramuscularly.

565 ^e Reduction in rate was adjusted for country, age, prior influenza vaccination status, and wheezing history status

566 A randomized, double-blind, saline placebo-controlled trial (D153-P501) was performed to evaluate the
567 efficacy of FluMist in children 12 through 35 months of age without high-risk medical conditions against
568 culture-confirmed influenza illness. This study was performed in Asia over two successive seasons (2000-
569 2001 and 2001-2002). The primary endpoint of the trial was the prevention of culture-confirmed influenza
570 illness due to antigenically matched wild-type influenza. Respiratory illness that prompted an influenza
571 culture was defined as at least one of the following: fever ($\geq 100.4^{\circ}\text{F}$ rectal or $\geq 99.5^{\circ}\text{F}$ axillary),
572 wheezing, shortness of breath, pulmonary congestion, pneumonia, or otitis media; or two of the following:
573 runny nose/nasal congestion, sore throat, cough, muscle aches, chills, headache, irritability, decreased
574 activity, or vomiting. A total of 3174 children were randomized 3:2 (vaccine: placebo) to receive 2 doses
575 of study vaccine or placebo at least 28 days apart in Year 1. See Table 7 for a description of the results.

576 During the second year of Study D153-P501, for children who received two doses in Year 1 and one dose
577 in Year 2, FluMist demonstrated 84.3% (95% CI: 70.1, 92.4) efficacy against culture-confirmed influenza
578 illness due to antigenically matched wild-type influenza.

579 Study AV006 was a second multi-center, randomized, double-blind, AF-SPG placebo-controlled trial
580 performed in U.S. children without high-risk medical conditions to evaluate the efficacy of FluMist against
581 culture-confirmed influenza over two successive seasons (1996-1997 and 1997-1998). The primary
582 endpoint of the trial was the prevention of culture-confirmed influenza illness due to antigenically matched
583 wild-type influenza in children who received two doses of vaccine in the first year and a single
584 revaccination dose in the second year. Respiratory illness that prompted an influenza culture was defined
585 as at least one of the following: fever ($\geq 101^{\circ}\text{F}$ rectal or oral; or $\geq 100.4^{\circ}\text{F}$ axillary), wheezing, shortness of
586 breath, pulmonary congestion, pneumonia, or otitis media; or two of the following: runny nose/nasal
587 congestion, sore throat, cough, muscle aches, chills, headache, irritability, decreased activity, or vomiting.
588 During the first year of the study, 1602 children 15 through 71 months of age were randomized 2:1
589 (vaccine: placebo). See Table 7 for a description of the results.

590
591

Table 7: Efficacy^a of FluMist vs. Placebo Against Culture-Confirmed Influenza Illness Due to Antigenically Matched Wild-Type Strains (Studies D153-P501^b & AV006^c, Year 1)

	D153-P501 ^d			AV006 ^e		
	FluMist n ^f (%)	Placebo n ^f (%)	% Efficacy (95% CI)	FluMist n ^f (%)	Placebo n ^f (%)	% Efficacy (95% CI)
	N^g = 1653	N^g = 1111		N^g = 849	N^g = 410	
Any strain	56 (3.4%)	139 (12.5%)	72.9% ^h (62.8, 80.5)	10 (1%)	73 (18%)	93.4% (87.5, 96.5)
A/H1N1	23 (1.4%)	81 (7.3%)	80.9% (69.4, 88.5) ⁱ	0	0	--
A/H3N2	4 (0.2%)	27 (2.4%)	90.0% (71.4, 97.5)	4 (0.5%)	48 (12%)	96.0% (89.4, 98.5)
B	29 (1.8%)	35 (3.2%)	44.3% (6.2, 67.2)	6 (0.7%)	31 (7%)	90.5% (78.0, 95.9)

592 ^a D153-P501 and AV006 data are for subjects who received two doses of study vaccine.
593 ^b In children 12 through 35 months of age
594 ^c In children 15 through 71 months of age
595 ^d NCT00192244; see www.clinicaltrials.gov
596 ^e NCT00192179; see www.clinicaltrials.gov
597 ^f Number and percent of subjects in per-protocol efficacy analysis population with culture-confirmed influenza illness.
598 ^g Number of subjects in per-protocol efficacy analysis population of each treatment group of each study for the “any
599 strain” analysis.
600 ^h For D153-P501, influenza circulated through 12 months following vaccination.
601 ⁱ Estimate includes A/H1N1 and A/H1N2 strains. Both were considered antigenically similar to the vaccine.

602 During the second year of Study AV006, children remained in the same treatment group as in Year 1 and
603 received a single dose of FluMist or placebo. During the second year, the primary circulating strain was
604 the A/Sydney/05/97 H3N2 strain, which was antigenically dissimilar from the H3N2 strain represented in
605 the vaccine, A/Wuhan/359/95; FluMist demonstrated 87.0% (95% CI: 77.0, 92.6) efficacy against culture-
606 confirmed influenza illness.

607 **14.2 Immune Response Study of FluMist Quadrivalent in Children and Adolescents**

608 A multicenter, randomized, double-blind, active-controlled, non-inferiority study (MI-CP208) was
609 performed to assess the immunogenicity of FluMist Quadrivalent compared to FluMist (active control) in
610 children and adolescents 2 through 17 years of age. A total of 2312 subjects were randomized by site at a
611 3:1:1 ratio to receive either FluMist Quadrivalent or one of two formulations of comparator vaccine
612 FluMist, each containing a B strain that corresponded to one of the two B strains in FluMist Quadrivalent
613 (a B strain of the Yamagata lineage or a B strain of the Victoria lineage).

614 Children 2 through 8 years of age received 2 doses of vaccine approximately 30 days apart; children
615 9 years of age and older received 1 dose. For children 2 through 8 years of age with a history of influenza
616 vaccination, immunogenicity assessments were performed prior to vaccination and at 28 days after the
617 first dose. For children 2 through 8 years of age without a history of influenza vaccination, immunogenicity
618 assessments were performed prior to vaccination and 28 days after the second dose. For children
619 9 years of age and older, immunogenicity assessments were performed prior to vaccination and at
620 28 days post vaccination.

621 Immunogenicity was evaluated by comparing the 4 strain-specific serum hemagglutination inhibition (HAI)
 622 antibody geometric mean titers (GMTs) post dosing and provided evidence that the addition of the second
 623 B strain did not result in immune interference to other strains included in the vaccine.

624 **14.3 Effectiveness Study of FluMist in Adults**

625 AV009 was a U.S. multi-center, randomized, double-blind, AF-SPG placebo-controlled trial to evaluate
 626 effectiveness of FluMist in adults 18 through 64 years of age without high-risk medical conditions over the
 627 1997-1998 influenza season. Participants were randomized 2:1 (vaccine: placebo). Cultures for influenza
 628 virus were not obtained from subjects in the trial, thus efficacy against culture-confirmed influenza was
 629 not assessed. The A/Wuhan/359/95 (H3N2) strain, which was contained in FluMist, was antigenically
 630 distinct from the predominant circulating strain of influenza virus during the trial period, A/Sydney/05/97
 631 (H3N2). Type A/Wuhan (H3N2) and Type B strains also circulated in the U.S. during the study period.
 632 The primary endpoint of the trial was the reduction in the proportion of participants with one or more
 633 episodes of any febrile illness, and prospective secondary endpoints were severe febrile illness and
 634 febrile upper respiratory illness. Effectiveness for any of the three endpoints was not demonstrated in a
 635 subgroup of adults 50 through 64 years of age. Primary and secondary effectiveness endpoints from the
 636 age group 18 through 49 years are presented in Table 8. Effectiveness was not demonstrated for the
 637 primary endpoint in adults 18 through 49 years of age.

638 **Table 8: Effectiveness of FluMist to Prevent Febrile Illness in Adults 18 through 49 Years of Age**
 639 **During the 7-Week Site-Specific Outbreak Period (Study AV009)**

Endpoint	FluMist N = 2411 ^a n (%)	Placebo N = 1226 ^a n (%)	Percent Reduction	(95% CI)
Participants with one or more events of:^b				
Primary Endpoint:				
Any febrile illness	331 (13.73)	189 (15.42)	10.9	(-5.1, 24.4)
Secondary Endpoints:				
Severe febrile illness	250 (10.37)	158 (12.89)	19.5	(3.0, 33.2)
Febrile upper respiratory illness	213 (8.83)	142 (11.58)	23.7	(6.7, 37.5)

640 ^a Number of evaluable subjects (92.7% and 93.0% of FluMist and placebo recipients, respectively).

641 ^b The predominantly circulating virus during the trial period was A/Sydney/05/97 (H3N2), an antigenic variant not
 642 included in the vaccine.

643 Effectiveness was shown in a post-hoc analysis using an endpoint of CDC-ILI in the age group 18
 644 through 49 years of age.

645 **14.4 Immune Response Study of FluMist Quadrivalent in Adults**

646 A multicenter, randomized, double-blind, active-controlled, and non-inferiority study (MI-CP185) was
 647 performed to assess the safety and immunogenicity of FluMist Quadrivalent compared to those of FluMist
 648 (active control) in adults 18 through 49 years of age. A total of 1800 subjects were randomized by site at
 649 a 4:1:1 ratio to receive either 1 dose of FluMist Quadrivalent or 1 dose of one of two formulations of

650 comparator vaccine, FluMist, each containing a B strain that corresponded to one of the two B strains in
651 FluMist Quadrivalent (a B strain of the Yamagata lineage and a B strain of the Victoria lineage).

652 Immunogenicity in study MI-CP185 was evaluated by comparing the 4 strain-specific serum
653 hemagglutination inhibition (HAI) antibody geometric mean titers (GMTs) post dosing and provided
654 evidence that the addition of the second B strain did not result in immune interference to other strains
655 included in the vaccine.

656 **14.5 Concomitantly Administered Live Virus Vaccines**

657 In Study AV018, concomitant administration of FluMist, MMR (manufactured by Merck & Co., Inc.) and
658 Varicella Virus Vaccine Live (manufactured by Merck & Co., Inc.) was studied in 1245 subjects 12
659 through 15 months of age. Subjects were randomized in a 1:1:1 ratio to MMR, Varicella vaccine and AF-
660 SPG placebo (group 1); MMR, Varicella vaccine and FluMist (group 2); or FluMist alone (group 3).
661 Immune responses to MMR and Varicella vaccines were evaluated 6 weeks post-vaccination while the
662 immune responses to FluMist were evaluated 4 weeks after the second dose. No evidence of
663 interference with immune response to measles, mumps, rubella, varicella and FluMist vaccines was
664 observed.

665 **15 REFERENCES**

666 1. Lasky T, Terracciano GJ, Magder L, et al. The Guillain-Barré syndrome and the 1992 – 1993 and
667 1993 – 1994 influenza vaccines. N Engl J Med 1998;339(25):1797-802.

668 **16 HOW SUPPLIED/STORAGE AND HANDLING**

669 **16.1 How Supplied**

670 FluMist Quadrivalent is supplied in a package of 10 pre-filled, single-dose (0.2 mL) intranasal sprayers.
671 The single-use intranasal sprayer is not made with natural rubber latex.
672 Carton containing 10 intranasal sprayers: NDC 66019-302-10
673 Single intranasal sprayer: NDC 66019-302-01

674 **16.2 Storage and Handling**

675 The cold chain [2-8°C (35-46°F)] must be maintained when transporting FluMist Quadrivalent.

676 **FLUMIST QUADRIVALENT SHOULD BE STORED IN A REFRIGERATOR BETWEEN 2-8°C (35-46°F)**
677 **UPON RECEIPT. THE PRODUCT MUST BE USED BEFORE THE EXPIRATION DATE ON THE**
678 **SPRAYER LABEL.**

679 **DO NOT FREEZE.**

680 Keep FluMist Quadrivalent sprayer in outer carton in order to protect from light.

681 A single temperature excursion up to 25°C (77°F) for 12 hours has been shown to have no adverse
682 impact on the vaccine. After a temperature excursion, the vaccine should be returned immediately to the
683 recommended storage condition (2°C – 8°C) and used as soon as feasible. Subsequent excursions are
684 not permitted.

685 Once FluMist Quadrivalent has been administered or has expired, the sprayer should be disposed of
686 according to the standard procedures for medical waste (e.g., sharps container or biohazard container).

687 **17 PATIENT COUNSELING INFORMATION**

688 **Advise the vaccine recipient or caregiver to read the FDA-approved patient labeling (Information**
689 **for Patients and Their Caregivers).**

690 Inform vaccine recipients or their parents/guardians of the need for two doses at least 1 month apart in
691 children 2 through 8 years of age, depending on vaccination history. Provide the Vaccine Information
692 Statements (VIS) which are required by the National Childhood Vaccine Injury Act of 1986 to be given
693 with each immunization.

694 **17.1 Asthma and Recurrent Wheezing**

695 Ask the vaccinee or their parent/guardian if the vaccinee has asthma. For children younger than 5 years
696 of age, also ask if the vaccinee has recurrent wheezing since this may be an asthma equivalent in this
697 age group. Inform the vaccinee or their parent/guardian that there may be an increased risk of wheezing
698 associated with FluMist Quadrivalent in persons younger than 5 years of age with recurrent wheezing and
699 persons of any age with asthma [see *Warnings and Precautions (5.2)*].

700 **17.2 Vaccination with a Live Virus Vaccine**

701 Inform vaccine recipients or their parents/guardians that FluMist Quadrivalent is an attenuated live virus
702 vaccine and has the potential for transmission to immunocompromised household contacts.

703 **17.3 Adverse Event Reporting**

704 Instruct the vaccine recipient or their parent/guardian to report adverse reactions to their healthcare
705 provider.

706 FluMist[®] is a registered trademark of MedImmune, LLC.

707  **MedImmune**

708 **Manufactured by:**

709 MedImmune, LLC

710 Gaithersburg, MD 20878

711 1-877-633-4411

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715 **Information for Patients and Their Caregivers**
716 **FluMist® Quadrivalent** (pronounced FLEW-mĭst Kwä-drē-VĀ-lənt)
717 (Influenza Vaccine Live, Intranasal)

718 **Please read this Patient Information carefully before you or your child is vaccinated with FluMist**
719 **Quadrivalent.**

720 This is a summary of information about FluMist Quadrivalent. It does not take the place of talking with
721 your healthcare provider about influenza vaccination. If you have questions or would like more
722 information, please talk with your healthcare provider.

723 **What is FluMist Quadrivalent?**

724 FluMist Quadrivalent is a vaccine that is sprayed into the nose to help protect against influenza. It can be
725 used in children, adolescents, and adults ages 2 through 49. FluMist Quadrivalent is similar to
726 MedImmune's trivalent Influenza Vaccine Live, Intranasal (FluMist) except FluMist Quadrivalent provides
727 protection against an additional influenza strain. FluMist Quadrivalent may not prevent influenza in
728 everyone who gets vaccinated.

729 **Who should not get FluMist Quadrivalent?**

730 You should not get FluMist Quadrivalent if you:

- 731 • have a severe allergy to eggs or to any inactive ingredient in the vaccine (see "What are the
732 ingredients in FluMist Quadrivalent?")
- 733 • have ever had a life-threatening reaction to influenza vaccinations
- 734 • are 2 through 17 years old and take aspirin or medicines containing aspirin. Children or
735 adolescents should not be given aspirin for 4 weeks after getting FluMist or FluMist
736 Quadrivalent unless your healthcare provider tells you otherwise.

737 Please talk to your healthcare provider if you are not sure if the items listed above apply to you or your
738 child.

739 Children under 2 years old have an increased risk of wheezing (difficulty with breathing) after getting
740 FluMist Quadrivalent.

741 **Who may not be able to get FluMist Quadrivalent?**

742 Tell your healthcare provider if you or your child:

- 743 • are currently wheezing
- 744 • have a history of wheezing if under 5 years old
- 745 • have had Guillain-Barré syndrome
- 746 • have a weakened immune system or live with someone who has a severely weakened
747 immune system

- 748 • have problems with your heart, kidneys, or lungs
- 749 • have diabetes
- 750 • are pregnant or nursing
- 751 • are taking Tamiflu[®], Relenza[®], amantadine, or rimantadine

752 If you or your child cannot take FluMist Quadrivalent, you may still be able to get an influenza shot. Talk
753 to your healthcare provider about this.

754 **How is FluMist Quadrivalent given?**

- 755 • FluMist Quadrivalent is a liquid that is sprayed into the nose.
- 756 • You can breathe normally while getting FluMist Quadrivalent. There is no need to inhale or
757 “sniff” it.
- 758 • People 9 years of age and older need one dose of FluMist Quadrivalent each year.
- 759 • Children 2 through 8 years old may need 2 doses of FluMist Quadrivalent, depending on their
760 history of previous influenza vaccination. Your healthcare provider will decide if your child
761 needs to come back for a second dose.

762 **What are the possible side effects of FluMist Quadrivalent?**

763 **The most common side effects** are:

- 764 • runny or stuffy nose
- 765 • sore throat
- 766 • fever over 100 degrees F

767 **Other possible side effects** include:

- 768 • decreased appetite
- 769 • irritability
- 770 • tiredness
- 771 • cough
- 772 • headache
- 773 • muscle ache
- 774 • chills

775 Call your healthcare provider or go to the emergency department right away if you or your child
776 experience:

- 777 • hives or a bad rash
- 778 • trouble breathing
- 779 • swelling of the face, tongue, or throat

780 These are not all the possible side effects of FluMist Quadrivalent. You can ask your healthcare provider
781 for a complete list of side effects that is available to healthcare professionals.

782 Call your healthcare provider for medical advice about side effects. You may report side effects to VAERS
783 at 1-800-822-7967 or <http://vaers.hhs.gov>.

784 **What are the ingredients in FluMist Quadrivalent?**

785 Active Ingredient: FluMist Quadrivalent contains 4 influenza virus strains that are weakened (A(H1N1),
786 A(H3N2), B Yamagata lineage, and B Victoria lineage).

787 Inactive Ingredients: monosodium glutamate, gelatin, arginine, sucrose, dibasic potassium phosphate,
788 monobasic potassium phosphate, and gentamicin.

789 FluMist Quadrivalent does not contain preservatives.

790 **How is FluMist Quadrivalent Stored?**

791 FluMist Quadrivalent is stored in a refrigerator (not the freezer) between 35-46 degrees F (2-8 degrees C)
792 upon receipt. FluMist Quadrivalent sprayer must be kept in the carton until use in order to protect from
793 light. FluMist Quadrivalent must be used before the expiration date on the sprayer label.

794 If you would like more information, talk to your healthcare provider or visit www.flumistquadrivalent.com or
795 call 1-877-633-4411.

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797 Other brands listed are registered trademarks of their respective owners and are not trademarks of
798 MedImmune, LLC.



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