

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

**HAVRIX (Hepatitis A Vaccine)
Suspension for Intramuscular Injection
Initial U.S. Approval: 1995**

INDICATIONS AND USAGE

HAVRIX is a vaccine indicated for active immunization against disease caused by hepatitis A virus (HAV). HAVRIX is approved for use in persons 12 months of age and older. Primary immunization should be administered at least 2 weeks prior to expected exposure to HAV. (1)

DOSAGE AND ADMINISTRATION

- HAVRIX is administered by intramuscular injection. (2.2)
- Children and adolescents: A single 0.5-mL dose and a 0.5-mL booster dose administered between 6 to 12 months later. (2.3)
- Adults: A single 1-mL dose and a 1-mL booster dose administered between 6 to 12 months later. (2.3)

DOSAGE FORMS AND STRENGTHS

- Suspension for injection available in the following presentations:
 - 0.5-mL single-dose vials and prefilled syringes. (3)
 - 1-mL single-dose vials and prefilled syringes. (3)

CONTRAINDICATIONS

Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any hepatitis A-containing vaccine, or to any component of HAVRIX, including neomycin. (4)

WARNINGS AND PRECAUTIONS

- The tip caps of the prefilled syringes may contain natural rubber latex which may cause allergic reactions in latex-sensitive individuals. (5.1)
- Syncope (fainting) can occur in association with administration of injectable vaccines, including HAVRIX. Procedures should be in place to avoid falling injury and to restore cerebral perfusion following syncope. (5.2)

ADVERSE REACTIONS

- In studies of adults and children 2 years of age and older, the most common solicited adverse events were injection-site soreness (56% of adults and 21% of children) and headache (14% of adults and less than 9% of children). (6.1)
- In studies of children 11 to 25 months of age, the most frequently reported solicited local reactions were pain (32%) and redness (29%). Common solicited general adverse events were irritability (42%), drowsiness (28%), and loss of appetite (28%). (6.1)

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

DRUG INTERACTIONS

Do not mix HAVRIX with any other vaccine or product in the same syringe or vial. (7.1)

USE IN SPECIFIC POPULATIONS

Safety and effectiveness of HAVRIX have not been established in pregnant women and nursing mothers. (8.1)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: XX/XXXX

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1 FULL PRESCRIBING INFORMATION

2 1 INDICATIONS AND USAGE

3 HAVRIX[®] is indicated for active immunization against disease caused by hepatitis A
4 virus (HAV). HAVRIX is approved for use in persons 12 months of age and older. Primary
5 immunization should be administered at least 2 weeks prior to expected exposure to HAV.

6 2 DOSAGE AND ADMINISTRATION

7 2.1 Preparation for Administration

8 Shake well before use. With thorough agitation, HAVRIX is a homogeneous, turbid,
9 white suspension. Do not administer if it appears otherwise. Parenteral drug products should be
10 inspected visually for particulate matter and discoloration prior to administration, whenever
11 solution and container permit. If either of these conditions exists, the vaccine should not be
12 administered.

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

14 For the vials, use a sterile needle and sterile syringe to withdraw the vaccine dose and
15 administer intramuscularly. Changing needles between drawing vaccine from a vial and injecting
16 it into a recipient is not necessary unless the needle has been damaged or contaminated. Use a
17 separate sterile needle and syringe for each individual.

18 2.2 Administration

19 HAVRIX should be administered by intramuscular injection only. HAVRIX should not
20 be administered in the gluteal region; such injections may result in suboptimal response.

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

22 2.3 Recommended Dose and Schedule

23 Children and Adolescents: Primary immunization for children and adolescents
24 (12 months through 18 years of age) consists of a single 0.5-mL dose and a 0.5-mL booster dose
25 administered anytime between 6 and 12 months later. The preferred sites for intramuscular
26 injections are the anterolateral aspect of the thigh in young children or the deltoid muscle of the
27 upper arm in older children.

28 Adults: Primary immunization for adults consists of a single 1-mL dose and a 1-mL
29 booster dose administered anytime between 6 and 12 months later. In adults, the injection should
30 be given in the deltoid region.

31 3 DOSAGE FORMS AND STRENGTHS

32 Suspension for injection available in the following presentations:

- 33 • 0.5-mL single-dose vials and prefilled TIP-LOK[®] syringes.
- 34 • 1-mL single-dose vials and prefilled TIP-LOK syringes.

35 **4 CONTRAINDICATIONS**

36 Severe allergic reaction (e.g., anaphylaxis) after a previous dose of any hepatitis A-
37 containing vaccine, or to any component of HAVRIX, including neomycin, is a contraindication
38 to administration of HAVRIX [*see Description (11)*].

39 **5 WARNINGS AND PRECAUTIONS**

40 **5.1 Latex**

41 The tip caps of the prefilled syringes may contain natural rubber latex which may cause
42 allergic reactions in latex-sensitive individuals.

43 **5.2 Syncope**

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

48 **5.3 Preventing and Managing Allergic Vaccine Reactions**

49 Appropriate medical treatment and supervision must be available to manage possible
50 anaphylactic reactions following administration of the vaccine [*see Contraindications (4)*].

51 **5.4 Altered Immunocompetence**

52 Immunocompromised persons may have a diminished immune response to HAVRIX,
53 including individuals receiving immunosuppressant therapy.

54 **5.5 Limitations of Vaccine Effectiveness**

55 Hepatitis A virus has a relatively long incubation period (15 to 50 days). HAVRIX may
56 not prevent hepatitis A infection in individuals who have an unrecognized hepatitis A infection at
57 the time of vaccination. Additionally, vaccination with HAVRIX may not protect all individuals.

58 **6 ADVERSE REACTIONS**

59 **6.1 Clinical Trials Experience**

60 Because clinical trials are conducted under widely varying conditions, adverse reaction
61 rates observed in the clinical trials of a vaccine cannot be directly compared to rates in the
62 clinical trials of another vaccine, and may not reflect the rates observed in practice.

63 The safety of HAVRIX has been evaluated in 61 clinical trials involving more than
64 34,000 individuals receiving doses of 360 EL.U., 720 EL.U., or 1440 EL.U.

65 Of solicited adverse events in clinical trials of adults, who received HAVRIX
66 1440 EL.U., and children (2 years of age and older), who received either HAVRIX 360 EL.U. or
67 720 EL.U., the most frequently reported was injection-site soreness (56% of adults and 21% of
68 children); less than 0.5% of soreness was reported as severe. Headache was reported by 14% of
69 adults and less than 9% of children. Other solicited and unsolicited events occurring during
70 clinical trials are listed below.

71 Incidence 1% to 10% of Injections: *Metabolism and Nutrition Disorders:* Anorexia.
72 *Gastrointestinal Disorders:* Nausea.

73 *General Disorders and Administration Site Conditions:* Fatigue, fever >99.5°F
74 (37.5°C), induration, redness, and swelling of the injection site; malaise.

75 Incidence <1% of Injections: *Infections and Infestations:* Pharyngitis, upper
76 respiratory tract infections.

77 *Blood and Lymphatic System Disorders:* Lymphadenopathy.

78 *Psychiatric Disorders:* Insomnia.

79 *Nervous System Disorders:* Dysgeusia, hypertonia.

80 *Eye Disorders:* Photophobia.

81 *Ear and Labyrinth Disorders:* Vertigo.

82 *Gastrointestinal Disorders:* Abdominal pain, diarrhea, vomiting.

83 *Skin and Subcutaneous Tissue Disorders:* Pruritus, rash, urticaria.

84 *Musculoskeletal and Connective Tissue Disorders:* Arthralgia, myalgia.

85 *General Disorders and Administration Site Conditions:* Injection site hematoma.

86 *Investigations:* Creatine phosphokinase increased.

87 Studies of HAVRIX 720 EL.U./0.5 mL in Children 11 to 25 Months of Age: In 4
88 studies, 3,152 children 11 to 25 months of age received at least one dose of HAVRIX 720 EL.U.
89 administered alone or concomitantly with other routine childhood vaccinations [*see Clinical*
90 *Studies (14.2, 14.5)*]. The studies included HAV 210 (N = 1,084), HAV 232 (N = 394),
91 HAV 220 (N = 433), and HAV 231 (N = 1,241).

92 In the largest of these studies (HAV 231) conducted in the US, 1,241 children 15 months
93 of age were randomized to receive: Group 1) HAVRIX alone; Group 2) HAVRIX concomitantly
94 with measles, mumps, and rubella (MMR) vaccine (manufactured by Merck and Co.) and
95 varicella vaccine (manufactured by Merck and Co.); or Group 3) MMR and varicella vaccines.
96 Subjects in Group 3 who received MMR and varicella vaccines received the first dose of
97 HAVRIX 42 days later. A second dose of HAVRIX was administered to all subjects 6 to
98 9 months after the first dose of HAVRIX. Solicited local adverse reactions and general events
99 were recorded by parents/guardians on diary cards for 4 days (days 0 to 3) after vaccination.
100 Unsolicited adverse events were recorded on the diary card for 31 days after vaccination.
101 Telephone follow-up was conducted 6 months after the last vaccination to inquire about serious
102 adverse events, new onset chronic illnesses and medically significant events. A total of 1,035
103 children completed the 6-month follow-up. Among subjects in all groups combined, 53% were
104 male; 69% of subjects were white, 16% were Hispanic, 9% were black and 6% were other
105 racial/ethnic groups.

106 Percentages of subjects with solicited local adverse reactions and general adverse events
107 following HAVRIX administered alone (Group 1) or concomitantly with MMR and varicella
108 vaccines (Group 2) are presented in Table 1. The solicited adverse events from the 3 additional
109 coadministration studies conducted with HAVRIX were comparable to those from Study
110 HAV 231.

111

112 **Table 1. Solicited Local Adverse Reactions and General Adverse Events Occurring Within**
 113 **4 Days of Vaccination^a in Children 15 to 24 Months of Age With HAVRIX Administered**
 114 **Alone or Concomitantly With MMR and Varicella Vaccines (TVC)**

	Group 1 HAVRIX Dose 1 %	Group 2 HAVRIX+ MMR+V^b Dose 1 %	Group 1 HAVRIX Dose 2 %	Group 2 HAVRIX Dose 2 %
Local (at injection site for HAVRIX)				
N	298	411	272	373
Pain, any	23.8	23.6	24.3	30.3
Redness, any	20.1	20.0	22.8	23.9
Swelling, any	8.7	10.2	9.6	9.9
General				
N	300	417	271	375
Irritability, any	33.3	43.9	31.0	27.2
Irritability, grade 3	0.3	1.9	1.5	0.3
Drowsiness, any	22.3	35.3	21.0	20.8
Drowsiness, grade 3	1.0	2.2	1.1	0.0
Loss of appetite, any	18.3	26.1	19.9	20.5
Loss of appetite, grade 3	1.0	1.4	0.4	0.3
Fever $\geq 100.6^{\circ}\text{F}$ (38.1°C)	3.0	4.8	3.3	2.7
Fever $\geq 101.5^{\circ}\text{F}$ (38.6°C)	2.0	2.6	1.8	1.6
Fever $\geq 102.4^{\circ}\text{F}$ (39.1°C)	0.7	0.7	0.4	1.1

115 Total vaccinated cohort (TVC) = all subjects who received at least one dose of vaccine.

116 N = number of subjects who received at least one dose of vaccine and for whom diary card information
 117 was available.

118 Grade 3: drowsiness defined as prevented normal daily activities; irritability/fussiness defined as crying
 119 that could not be comforted/prevented normal daily activities; loss of appetite defined as no eating at
 120 all.

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

122 ^b MMR = measles, mumps, and rubella vaccine; V = varicella vaccine.

123
 124 *Serious Adverse Events in Children 11 to 25 Months of Age:* Among these 4
 125 studies, 0.9% (29/3,152) of subjects reported a serious adverse event within the 31-day period
 126 following vaccination with HAVRIX. Among subjects administered HAVRIX alone 1.0%
 127 (13/1,332) reported a serious adverse event. Among subjects who received HAVRIX
 128 concomitantly with other childhood vaccines, 0.9% (8/909) reported a serious adverse event. In
 129 these 4 studies, there were 4 reports of seizure within 31 days post-vaccination: these occurred 2,
 130 9, and 27 days following the first dose of HAVRIX administered alone and 12 days following

131 the second dose of HAVRIX. In one subject who received INFANRIX and Hib conjugate
132 vaccine followed by HAVRIX 6 weeks later, bronchial hyperreactivity and respiratory distress
133 were reported on the day of administration of HAVRIX alone.

134 **6.2 Postmarketing Experience**

135 In addition to reports in clinical trials, worldwide voluntary reports of adverse events
136 received for HAVRIX since market introduction of this vaccine are listed below. This list
137 includes serious adverse events or events which have a suspected causal connection to
138 components of HAVRIX or other vaccines or drugs. Because these events are reported
139 voluntarily from a population of uncertain size, it is not always possible to reliably estimate their
140 frequency or establish a causal relationship to the vaccine.

141 Infections and Infestations: Rhinitis.

142 Blood and Lymphatic System Disorders: Thrombocytopenia.

143 Immune System Disorders: Anaphylactic reaction, anaphylactoid reaction, serum
144 sickness-like syndrome.

145 Nervous System Disorders: Convulsion, dizziness, encephalopathy, Guillain-Barré
146 syndrome, hypoesthesia, multiple sclerosis, myelitis, neuropathy, paresthesia, somnolence,
147 syncope.

148 Vascular Disorders: Vasculitis.

149 Respiratory, Thoracic, and Mediastinal Disorders: Dyspnea.

150 Hepatobiliary Disorders: Hepatitis, jaundice.

151 Skin and Subcutaneous Tissue Disorders: Angioedema, erythema multiforme,
152 hyperhidrosis.

153 Congenital, Familial, and Genetic Disorders: Congenital anomaly.

154 Musculoskeletal and Connective Tissue Disorders: Musculoskeletal stiffness.

155 General Disorders and Administration Site Conditions: Chills, influenza-like
156 symptoms, injection site reaction, local swelling.

157 **7 DRUG INTERACTIONS**

158 **7.1 Concomitant Administration With Vaccines and Immune Globulin**

159 In clinical studies HAVRIX was administered concomitantly with the following vaccines
160 [*see Adverse Reactions (6.1) and Clinical Studies (14.5)*]:

- 161 • INFANRIX (DTaP);
- 162 • Hib conjugate vaccine;
- 163 • pneumococcal 7-valent conjugate vaccine;
- 164 • MMR vaccine;
- 165 • varicella vaccine.

166 HAVRIX may be administered concomitantly with immune globulin.

167 When concomitant administration of other vaccines or immune globulin is required, they
168 should be given with different syringes and at different injection sites. Do not mix HAVRIX with
169 any other vaccine or product in the same syringe or vial.

170 **7.2 Immunosuppressive Therapies**

171 Immunosuppressive therapies, including irradiation, antimetabolites, alkylating agents,
172 cytotoxic drugs, and corticosteroids (used in greater than physiologic doses), may reduce the
173 immune response to HAVRIX.

174 **8 USE IN SPECIFIC POPULATIONS**

175 **8.1 Pregnancy**

176 Pregnancy Category C

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

181 **8.3 Nursing Mothers**

182 It is not known whether HAVRIX is excreted in human milk. Because many drugs are
183 excreted in human milk, caution should be exercised when HAVRIX is administered to a nursing
184 woman.

185 **8.4 Pediatric Use**

186 The safety and effectiveness of HAVRIX, doses of 360 EL.U. or 720 EL.U., have been
187 evaluated in more than 22,000 subjects 1 year to 18 years of age.

188 The safety and effectiveness of HAVRIX have not been established in subjects younger
189 than 12 months of age.

190 **8.5 Geriatric Use**

191 Clinical studies of HAVRIX did not include sufficient numbers of subjects 65 years of
192 age and older to determine whether they respond differently from younger subjects. Other
193 reported clinical experience has not identified differences in overall safety between these
194 subjects and younger adult subjects.

195 **8.6 Hepatic Impairment**

196 Subjects with chronic liver disease had a lower antibody response to HAVRIX than
197 healthy subjects [*see Clinical Studies (14.3)*].

198 **11 DESCRIPTION**

199 HAVRIX (Hepatitis A Vaccine) is a sterile suspension of inactivated virus for
200 intramuscular administration. The virus (strain HM175) is propagated in MRC-5 human diploid
201 cells. After removal of the cell culture medium, the cells are lysed to form a suspension. This
202 suspension is purified through ultrafiltration and gel permeation chromatography procedures.
203 Treatment of this lysate with formalin ensures viral inactivation. Viral antigen activity is
204 referenced to a standard using an enzyme linked immunosorbent assay (ELISA), and is therefore
205 expressed in terms of ELISA Units (EL.U.).

206 Each 1-mL adult dose of vaccine contains 1440 EL.U. of viral antigen, adsorbed on
207 0.5 mg of aluminum as aluminum hydroxide.

208 Each 0.5-mL pediatric dose of vaccine contains 720 EL.U. of viral antigen, adsorbed onto
209 0.25 mg of aluminum as aluminum hydroxide.

210 HAVRIX contains the following excipients: Amino acid supplement (0.3% w/v) in a
211 phosphate-buffered saline solution and polysorbate 20 (0.05 mg/mL). From the manufacturing
212 process, HAVRIX also contains residual MRC-5 cellular proteins (not more than 5 mcg/mL),
213 formalin (not more than 0.1 mg/mL), and neomycin sulfate (not more than 40 ng/mL), an
214 aminoglycoside antibiotic included in the cell growth media.

215 HAVRIX is formulated without preservatives.

216 HAVRIX is available in vials and prefilled syringes. The tip caps of the prefilled syringes
217 may contain natural rubber latex; the plungers are not made with natural rubber latex. The vial
218 stoppers are not made with natural rubber latex.

219 **12 CLINICAL PHARMACOLOGY**

220 **12.1 Mechanism of Action**

221 The hepatitis A virus belongs to the picornavirus family. It is one of several hepatitis
222 viruses that cause systemic disease with pathology in the liver.

223 The incubation period for hepatitis A averages 28 days (range: 15 to 50 days).¹ The
224 course of hepatitis A infection is extremely variable, ranging from asymptomatic infection to
225 icteric hepatitis and death.

226 The presence of antibodies to HAV confers protection against hepatitis A infection.
227 However, the lowest titer needed to confer protection has not been determined.

228 **13 NONCLINICAL TOXICOLOGY**

229 **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

230 HAVRIX has not been evaluated for its carcinogenic potential, mutagenic potential, or
231 potential for impairment of fertility.

232 **14 CLINICAL STUDIES**

233 **14.1 Pediatric Effectiveness Studies**

234 Protective efficacy with HAVRIX has been demonstrated in a double-blind, randomized
235 controlled study in school children (age 1 to 16 years) in Thailand who were at high risk of HAV
236 infection. A total of 40,119 children were randomized to be vaccinated with either HAVRIX
237 360 EL.U. or ENGERIX-B 10 mcg at 0, 1, and 12 months. Of these, 19,037 children received 2
238 doses of HAVRIX (0 and 1 months) and 19,120 children received 2 doses of control vaccine,
239 ENGERIX-B (0 and 1 months). A total of 38,157 children entered surveillance at day 138 and
240 were observed for an additional 8 months. Using the protocol-defined endpoint (≥ 2 days absence
241 from school, ALT level >45 U/mL, and a positive result in the HAVAB-M test), 32 cases of
242 clinical hepatitis A occurred in the control group. In the HAVRIX group, 2 cases were identified.
243 These 2 cases were mild in terms of both biochemical and clinical indices of hepatitis A disease.
244 Thus the calculated efficacy rate for prevention of clinical hepatitis A was 94% (95% Confidence
245 Interval [CI]: 74, 98).

246 In outbreak investigations occurring in the trial, 26 clinical cases of hepatitis A (of a total
247 of 34 occurring in the trial) occurred. No cases occurred in vaccinees who received HAVRIX.

248 Using additional virological and serological analyses post hoc, the efficacy of HAVRIX
249 was confirmed. Up to 3 additional cases of mild clinical illness may have occurred in vaccinees.
250 Using available testing, these illnesses could neither be proven nor disproven to have been
251 caused by HAV. By including these as cases, the calculated efficacy rate for prevention of
252 clinical hepatitis A would be 84% (95% CI: 60, 94).

253 **14.2 Immunogenicity in Children and Adolescents**

254 Immune Response to HAVRIX 720 EL.U./0.5 mL at 11 to 25 Months of Age

255 (Study HAV 210): In this prospective, open-label, multicenter study, 1,084 children were
256 administered study vaccine in one of 5 groups:

257 (1) Children 11 to 13 months of age who received HAVRIX on a 0- and 6-month
258 schedule;

259 (2) Children 15 to 18 months of age who received HAVRIX on a 0- and 6-month
260 schedule;

261 (3) Children 15 to 18 months of age who received HAVRIX coadministered with
262 INFANRIX and Haemophilus b (Hib) conjugate vaccine (no longer US-licensed) at month 0 and
263 HAVRIX at month 6;

264 (4) Children 15 to 18 months of age who received INFANRIX coadministered with Hib
265 conjugate vaccine at month 0 and HAVRIX at months 1 and 7;

266 (5) Children 23 to 25 months of age who received HAVRIX on a 0- and 6-month
267 schedule.

268 Among subjects in all groups, 52% were male; 61% of subjects were white, 9% were
269 black, 3% were Asian, and 27% were other racial/ethnic groups. The anti-hepatitis A antibody
270 vaccine responses and GMTs, calculated on responders for groups 1, 2, and 5 are presented in
271 Table 2. Vaccine response rates were similar among the 3 age groups that received HAVRIX.
272 One month after the second dose of HAVRIX, the GMT in each of the younger age groups (11 to
273 13 and 15 to 18 months of age) was shown to be similar to that achieved in the 23 to 25 months
274 of age group.

275

276 **Table 2. Anti-Hepatitis A Immune Response Following 2 Doses of HAVRIX**
 277 **720 EL.U./0.5 mL Administered 6 Months Apart in Children Given the First Dose of**
 278 **HAVRIX at 11 to 13 Months of Age, 15 to 18 Months of Age, or 23 to 25 Months of Age**

Age group	N	Vaccine Response		GMT (mIU/mL)
		%	95% CI	
11-13 months (Group 1)	218	99	97, 100	1,461 ^a
15-18 months (Group 2)	200	100	98, 100	1,635 ^a
23-25 months (Group 5)	211	100	98, 100	1,911

279 Vaccine response = Seroconversion (anti-HAV ≥ 15 mIU/mL [lower limit of antibody
 280 measurement by assay]) in children initially seronegative or at least the maintenance of the
 281 pre-vaccination anti-HAV concentration in initially seropositive children.

282 CI = Confidence Interval; GMT = Geometric mean antibody titer.

283 ^a Calculated on vaccine responders one month post-dose 2. GMTs in children 11 to 13 months
 284 of age and 15 to 18 months of age were non-inferior (similar) to the GMT in children 23 to
 285 25 months of age (i.e., the lower limit of the two-sided 95% CI on the GMT ratio for
 286 Group 1/Group 5 and for Group 2/Group 5 were both ≥ 0.5).

287
 288 In 3 additional clinical studies (HAV 232, HAV 220, and HAV 231), children received
 289 either 2 doses of HAVRIX alone or the first dose of HAVRIX concomitantly administered with
 290 other routinely recommended US-licensed vaccines followed by a second dose of HAVRIX.
 291 After the second dose of HAVRIX, there was no evidence for interference with the anti-HAV
 292 response in the children who received concomitantly administered vaccines compared to those
 293 who received HAVRIX alone. [See Adverse Reactions (6.1) and Clinical Studies (14.5).]

294 **Immune Response to HAVRIX 360 EL.U. Among Individuals 2 to 18 Years of**
 295 **Age:** In 6 clinical studies, 762 subjects 2 to 18 years of age received 2 doses of HAVRIX
 296 (360 EL.U.) given 1 month apart (GMT ranged from 197 to 660 mIU/mL). Ninety-nine percent
 297 of subjects seroconverted following 2 doses. When a third dose of HAVRIX 360 EL.U. was
 298 administered 6 months following the initial dose, all subjects were seropositive (anti-HAV
 299 ≥ 20 mIU/mL) 1 month following the third dose, with GMTs rising to a range of 3,388 to
 300 4,643 mIU/mL. In 1 study in which children were followed for an additional 6 months, all
 301 subjects remained seropositive.

302 **Immune Response to HAVRIX 720 EL.U./0.5 mL Among Individuals 2 to 19**
 303 **Years of Age:** In 4 clinical studies, 314 children and adolescents ranging from 2 to 19 years of
 304 age were immunized with 2 doses of HAVRIX 720 EL.U./0.5 mL given 6 months apart. One
 305 month after the first dose, seroconversion (anti-HAV ≥ 20 mIU/mL [lower limit of antibody
 306 measurement by assay]) ranged from 96.8% to 100%, with GMTs of 194 mIU/mL to
 307 305 mIU/mL. In studies in which sera were obtained 2 weeks following the initial dose,
 308 seroconversion ranged from 91.6% to 96.1%. One month following the booster dose at month 6,
 309 all subjects were seropositive, with GMTs ranging from 2,495 mIU/mL to 3,644 mIU/mL.

310 In an additional study in which the booster dose was delayed until 1 year following the
311 initial dose, 95.2% of the subjects were seropositive just prior to administration of the booster
312 dose. One month later, all subjects were seropositive, with a GMT of 2,657 mIU/mL.

313 **14.3 Immunogenicity in Adults**

314 More than 400 healthy adults 18 to 50 years of age in 3 clinical studies were given a
315 single 1440 EL.U. dose of HAVRIX. All subjects were seronegative for hepatitis A antibodies at
316 baseline. Specific humoral antibodies against HAV were elicited in more than 96% of subjects
317 when measured 1 month after vaccination. By day 15, 80% to 98% of vaccinees had already
318 seroconverted (anti-HAV ≥ 20 mIU/mL [lower limit of antibody measurement by assay]). GMTs
319 of seroconverters ranged from 264 to 339 mIU/mL at day 15 and increased to a range of 335 to
320 637 mIU/mL by 1 month following vaccination.

321 The GMTs obtained following a single dose of HAVRIX are at least several times higher
322 than that expected following receipt of immune globulin.

323 In a clinical study using 2.5 to 5 times the standard dose of immune globulin (standard
324 dose = 0.02 to 0.06 mL/kg), the GMT in recipients was 146 mIU/mL at 5 days
325 post-administration, 77 mIU/mL at month 1, and 63 mIU/mL at month 2.

326 In 2 clinical trials in which a booster dose of 1440 EL.U. was given 6 months following
327 the initial dose, 100% of vaccinees (n = 269) were seropositive 1 month after the booster dose,
328 with GMTs ranging from 3,318 mIU/mL to 5,925 mIU/mL. The titers obtained from this
329 additional dose approximate those observed several years after natural infection.

330 In a subset of vaccinees (n = 89), a single dose of HAVRIX 1440 EL.U. elicited specific
331 anti-HAV neutralizing antibodies in more than 94% of vaccinees when measured 1 month after
332 vaccination. These neutralizing antibodies persisted until month 6. One hundred percent of
333 vaccinees had neutralizing antibodies when measured 1 month after a booster dose given at
334 month 6.

335 Immunogenicity of HAVRIX was studied in subjects with chronic liver disease of
336 various etiologies. 189 healthy adults and 220 adults with either chronic hepatitis B (n = 46),
337 chronic hepatitis C (n = 104), or moderate chronic liver disease of other etiology (n = 70) were
338 vaccinated with HAVRIX 1440 EL.U. on a 0- and 6-month schedule. The last group consisted of
339 alcoholic cirrhosis (n = 17), autoimmune hepatitis (n = 10), chronic hepatitis/cryptogenic
340 cirrhosis (n = 9), hemochromatosis (n = 2), primary biliary cirrhosis (n = 15), primary sclerosing
341 cholangitis (n = 4), and unspecified (n = 13). At each time point, geometric mean antibody titers
342 (GMTs) were lower for subjects with chronic liver disease than for healthy subjects. At month 7,
343 the GMTs ranged from 478 mIU/mL (chronic hepatitis C) to 1,245 mIU/mL (healthy). One
344 month after the first dose, seroconversion rates in adults with chronic liver disease were lower
345 than in healthy adults. However, 1 month after the booster dose at month 6, seroconversion rates
346 were similar in all groups; rates ranged from 94.7% to 98.1%. The relevance of these data to the
347 duration of protection afforded by HAVRIX is unknown.

348 In subjects with chronic liver disease, local injection site reactions with HAVRIX were
349 similar among all 4 groups, and no serious adverse events attributed to the vaccine were reported
350 in subjects with chronic liver disease.

351 **14.4 Duration of Immunity**

352 The duration of immunity following a complete schedule of immunization with HAVRIX
353 has not been established.

354 **14.5 Immune Response to Concomitantly Administered Vaccines**

355 In 3 clinical studies HAVRIX was administered concomitantly with other routinely
356 recommended US-licensed vaccines: Study HAV 232: Diphtheria and tetanus toxoids and
357 acellular pertussis vaccine adsorbed (INFANRIX, DTaP) and Haemophilus b (Hib) conjugate
358 vaccine (tetanus toxoid conjugate) (manufactured by sanofi pasteur SA); Study HAV 220:
359 Pneumococcal 7-valent conjugate vaccine (PCV-7) (manufactured by Pfizer), and Study
360 HAV 231: MMR and varicella vaccines. [See Adverse Reactions (6.1).]

361 Concomitant Administration With DTaP and Hib Conjugate Vaccine (Study

362 HAV 232): In this US multicenter study, 468 subjects, children 15 months of age were
363 randomized to receive: Group 1) HAVRIX coadministered with INFANRIX and Hib conjugate
364 vaccine (n = 127); Group 2) INFANRIX and Hib conjugate vaccine alone followed by a first
365 dose of HAVRIX one month later (n = 132); or Group 3) HAVRIX alone (n = 135). All subjects
366 received a second dose of HAVRIX alone 6 to 9 months following the first dose. Among
367 subjects in all groups combined, 53% were male; 64% of subjects were white, 12% were black,
368 6% were Hispanic, and 18% were other racial/ethnic groups.

369 There was no evidence for reduced antibody response to diphtheria and tetanus toxoids
370 (percentage of subjects with antibody levels ≥ 0.1 mIU/mL to each antigen), pertussis antigens
371 (percentage of subjects with seroresponse, antibody concentrations ≥ 5 EL.U./mL in seronegative
372 subjects or post-vaccination antibody concentration ≥ 2 times the pre-vaccination antibody
373 concentration in seropositive subjects, and GMTs), or Hib (percentage of subjects with antibody
374 levels ≥ 1 mcg/mL to polyribosyl-ribitol phosphate, PRP) when HAVRIX was administered
375 concomitantly with INFANRIX and Hib conjugate vaccine (Group 1) relative to INFANRIX and
376 Hib conjugate vaccine administered together (Group 2).

377 Concomitant Administration With Pneumococcal 7-Valent Conjugate Vaccine

378 (Study HAV 220): In this US multicenter study, 433 children 15 months of age were
379 randomized to receive: Group 1) HAVRIX coadministered with PCV-7 vaccine (n = 137); Group
380 2) HAVRIX administered alone (n = 147); or Group 3) PCV-7 vaccine administered alone
381 (n = 149) followed by a first dose of HAVRIX one month later. All subjects received a second
382 dose of HAVRIX 6 to 9 months after the first dose. Among subjects in all groups combined,
383 53% were female; 61% of subjects were white, 16% were Hispanic, 15% were black, and 8%
384 were other racial/ethnic groups.

385 There was no evidence for reduced antibody response to PCV-7 (GMC to each serotype)
386 when HAVRIX was administered concomitantly with PCV-7 vaccine (Group 1) relative to PCV-
387 7 administered alone (Group 3).

388 Concomitant Administration With MMR and Varicella Vaccines (Study HAV 231):
389 In a US multicenter study, there was no evidence for interference in the immune response to
390 MMR and varicella vaccines (the percentage of subjects with pre-specified
391 seroconversion/seroresponse levels) administered at 15 months of age concomitantly with
392 HAVRIX relative to the response when MMR and varicella vaccines are administered without
393 HAVRIX. [See Adverse Reactions (6.1).]

394 **15 REFERENCES**

395 1. Centers for Disease Control and Prevention. Prevention of hepatitis A through active or
396 passive immunization: Recommendations of the Immunization Practices Advisory
397 Committee (ACIP). *MMWR* 2006;55(RR-7):1-23.

398 **16 HOW SUPPLIED/STORAGE AND HANDLING**

399 HAVRIX is available in single-dose vials and prefilled disposable TIP-LOK syringes
400 (packaged without needles) (Preservative Free Formulation):

401 720 EL.U./0.5 mL

402 NDC 58160-825-01 Vial in Package of 10: NDC 58160-825-11

403 NDC 58160-825-43 Syringe in Package of 10: NDC 58160-825-52

404 1440 EL.U./mL

405 NDC 58160-826-01 Vial in Package of 10: NDC 58160-826-11

406 NDC 58160-826-05 Syringe in Package of 1: NDC 58160-826-34

407 NDC 58160-826-43 Syringe in Package of 10: NDC 58160-826-52

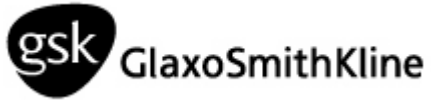
408 Store refrigerated between 2° and 8°C (36° and 46°F). Do not freeze. Discard if the
409 vaccine has been frozen. Do not dilute to administer.

410 **17 PATIENT COUNSELING INFORMATION**

- 411 • Inform vaccine recipients and parents or guardians of the potential benefits and risks of
412 immunization with HAVRIX.
- 413 • Emphasize, when educating vaccine recipients and parents or guardians regarding potential
414 side effects, that HAVRIX contains non-infectious killed viruses and cannot cause hepatitis
415 A infection.
- 416 • Instruct vaccine recipients and parents or guardians to report any adverse events to their
417 healthcare provider.
- 418 • Give vaccine recipients and parents or guardians the Vaccine Information Statements, which
419 are required by the National Childhood Vaccine Injury Act of 1986 to be given prior to
420 immunization. These materials are available free of charge at the Centers for Disease Control
421 and Prevention (CDC) website (www.cdc.gov/vaccines).

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