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TW:L1 PRESCRIBING INFORMATION	
Twinrix <sup>a</sup>	
Hepatitis A Inactivated & Hepatitis B	
(Recombinant) Vaccine DESCRIPTION	
Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] is a sterile bivalent vaccine containing the antigenic components used in producing Havrix® (Hepatitis A Vaccine, Inactivated) and Engerix-B® [Hepatitis B Vaccine (Recombinant)]. <i>Twinrix</i> is a sterile suspension of inactivated hepatitis A virus (strain HM175) propagated in MRC-5 cells, and combined with purified surface antigen of the hepatitis B virus. The purified hepatitis B surface antigen (HBsAg) is obtained by culturing genetically engineered <i>Saccharomyces cerevisiae</i> cells, which carry the surface antigen gene of the hepatitis B virus, in synthetic media containing inorganic salts, amino acids, dextrose, and vitamins. Bulk preparations of each antigen are adsorbed separately onto aluminum salts and then pooled during formulation.	
A 1.0 mL dose of vaccine contains not less than 720 ELISA Units of inactivated hepatitis A virus and 20 mcg of recombinant HBsAg protein.  One dose of vaccine also contains 0.45 mg of aluminum in the form of aluminum phosphate and aluminum hydroxide as adjuvants, amino acids, 5.0 mg 2-phenoxyethanol as a preservative, sodium chloride, phosphate buffer, polysorbate 20, Water for Injection, traces of formalin (not more than 0.1 mg), a trace amount of thimerosal (<1 mcg mercury) from the manufacturing process, and residual MRC-5 cellular proteins (not more than 2.5 mcg). Neomycin sulfate, an aminoglycoside antibiotic, is included in the cell growth media; only trace amounts (not more than 20 ng/dose) remain following purification. The manufacturing procedures used to manufacture <i>Twinrix</i> result in a product that contains no more than 5% yeast protein.	

Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] is supplied as a sterile suspension for intramuscular administration. The vaccine must be shaken before administration to ensure a uniform suspension. After shaking, the vaccine is a homogenous white turbid suspension.	
CLINICAL PHARMACOLOGY	
Several hepatitis viruses (A,B,C,D,E) are known to cause a systemic infection resulting in major pathologic changes in the liver. Features of hepatitis A and B are described below.	
Hepatitis A: The hepatitis A virus (HAV) belongs to the picornavirus family. Only one serotype of HAV has been described. <sup>1</sup>	
Hepatitis A is a highly contagious disease with the predominant mode of transmission being person-to-person via the fecal-oral route. Infection has been shown to be spread (1) by contaminated water or food; (2) by infected food handlers <sup>2</sup> ; (3) after breakdown in usual sanitary conditions or after floods or natural disasters; (4) by ingestion of raw or undercooked shellfish (oysters, clams, mussels) from contaminated waters <sup>3</sup> ; (5) during travel to areas of the world with poor hygienic conditions <sup>4</sup> ;(6) among institutionalized children and adults <sup>5</sup> ; (7) in daycare centers <sup>6</sup> ; (8) by parenteral transmission, either blood transfusions or sharing needles with infected people. <sup>7</sup>	
In the United States, attack rates for hepatitis A disease are cyclical and vary by population. The rates have increased gradually from 10.4 per 100,000 in 1987 to 11.7 per 100,000 in 1996.8	
The incubation period for hepatitis A averages 28 days (range: 15 to 50 days). The course of hepatitis A infection is extremely variable, ranging from asymptomatic infection to icteric hepatitis. However, most adults (76% to 97%) become symptomatic. Symptoms range from mild and transient to severe and prolonged and may include fever, nausea, vomiting and	

diarrhea in the prodromal phase, followed by jaundice in up to 88% of adults, as well as hepatomegaly and biochemical evidence of hepatocellular damage. Recovery is generally complete and followed by protection against HAV infection. However, illness may be prolonged, and relapse of clinical illness and viral shedding have been described. Up to 22% of adults who contract hepatitis A are hospitalized and approximately 100 patients die annually in the United States from complications of hepatitis A. 12	
Chronic shedding of HAV in feces has not been demonstrated, but relapses of hepatitis A can occur in as many as 20% of patients <sup>11,13</sup> and fecal shedding of HAV may recur at this time. <sup>11</sup> Approximately 70% of pediatric patients less than 6 years of age infected with hepatitis A are asymptomatic, and serve as a reservoir for infection among adults. <sup>12</sup>	
The presence of antibodies to HAV, as detected in a standardized assay (HAVAB), is an indication of the presence of protective antibodies against hepatitis A disease. Natural infection provides lifelong immunity even when antibodies to hepatitis A are undetectable. At present, studies show the duration of protection afforded by Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] against hepatitis A lasts at least 4 years. <sup>14</sup>	
Hepatitis B: The hepatitis B virus (HBV) belongs to a family of genetically related DNA-containing animal viruses, which are hepatotropic. The incubation period of hepatitis B ranges between 30 and 180 days. The mode of transmission of hepatitis B may be: by contact (contaminated body secretions including semen, vaginal secretions, blood, saliva); percutaneously (usually through accidental needlesticks or by sharing needles with infected people); or by maternal-neonatal transmission. <sup>15</sup>	
HBV infection occurs throughout the world with highly variable prevalences. A human reservoir of persistently infected persons is present in nearly all communities of the world. In the	

United States, parenteral drug abuse, unprotected sexual activity, occupationally acquired infection or travelers returning from high prevalence countries may be the principal mechanisms of HBV transmission.	
Clinical infection with hepatitis B may occur in two major forms: asymptomatic or symptomatic hepatitis. Asymptomatic HBV infection can be subclinical or inapparent. In subclinical infection, patients have abnormal liver enzymes without jaundice, while inapparent asymptomatic infection is identified only by serological testing. One in four adults who has symptomatic disease has jaundice (anicteric/icteric hepatitis).	
HBV infection can have serious consequences including acute massive hepatic necrosis, chronic active hepatitis and cirrhosis of the liver. As many as 90% of infants and 6% to 10% of adults who are infected in the United States will become HBV carriers. An estimated 200 to 300 million people are chronic carriers of HBV worldwide. The Centers for Disease Control and Prevention (CDC) estimates that there are approximately 1 million to 1.25 million chronic carriers of HBV in the United States. About 50,000 cases of hepatitis are reported per year, about half of which are hepatitis B. Unreported cases may be 10 times greater. Close contact (sexual contact or household contact) or exposure to blood from infected individuals is associated with increased risk of infection. Those patients who become chronic carriers can infect others and are at increased risk of developing primary hepatocellular carcinoma. Among other factors, infection with HBV may be the single most important factor for development of this carcinoma.	
Reduced Risk of Hepatocellular Carcinoma: A clear link has been demonstrated between chronic HBV infection and the occurrence of hepatocellular carcinoma. In a Taiwanese study, the institution of universal childhood immunization against HBV has been shown to decrease the incidence of hepatocellular carcinoma among children. <sup>17</sup> In a Korean study	

in adult males, vaccination against HBV has been shown to decrease the incidence of, and risk of, developing hepatocellular carcinoma in adults. <sup>18</sup>	
There is no definitive treatment for acute HBV infection. However, those who develop antibodies to HBsAg after active infection are protected against subsequent infection.  Antibody titers ≥10 mIU/mL against HBsAg are recognized as conferring protection against HBV. Seroconversion is defined as an antibody titer ≥1 mIU/mL.	
Clinical Trials	
Immunogenicity in Adults: Sera from 1,551 healthy adult volunteers ages 17 to 70, including 555 male subjects and 996 female subjects, in eleven clinical trials were analyzed following administration of three doses of Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] on a 0-, 1- and 6- month schedule. Seroconversion for antibodies against HAV was elicited in 99.9% of vaccinees, and protective antibodies against HBV were detected in 98.5%, one month after completion of the three-dose series.	

Table 1. Immunogenicity in *Twinrix* Worldwide Clinical Trials

Twinrix Dose	N	% Seroconversion for Hepatitis A*		% Seroprotection for Hepatitis B <sup>†</sup>
1	1,587	93.8		30.8
2	1,571	98.8		78.2
3	1,551	99.9		98.5
* anti-HAV titer ≥assay cut-off: 20 mIU/mL or 33 mIU/mL (Enzymun-Test®).  † anti-HBsAg titer ≥10 mIU/mL (AUSAB®).				
One of the eleven trials was a comparative trial conducted in a U.S. population given either <i>Twinrix</i> (on a 0-, 1-, 6-month schedule) or <i>Havrix</i> (0-, 6-month schedule) and <i>Engerix-B</i> (0-, 1-, 6-month schedule). The monovalent vaccines were given concurrently in opposite arms. Of a total of 773 adults (ages 18 to 70 years) enrolled in this trial, an immunogenicity analysis was performed in 533 subjects who				

completed the study according to protocol. Of these, 264 subjects received Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] and 269 subjects received Havrix and Engerix-B. Seroconversion against HAV and seroprotection against HBV are shown in Table 2.

Table 2. Percentage of Seroconversion or Seroprotection Rates in the *Twinrix* U.S. Clinical Trial

Vaccine	N	Time-point	% Seroconversion for Hepatitis A* (95% CI)	% Seroprotection for Hepatitis B <sup>†</sup> (95% CI)
Twinrix	264	Month 1	91.6	17.9
		Month 2	97.7	61.2
		Month 7	99.6 (97.9-100.0)	95.1 (91.7-97.4)
Havrix and Engerix-B	269	Month 1	98.1	7.5
9		Month 2	98.9	50.4
		Month 7	99.3 (97.3-99.9)	92.2 (88.3-95.1)
* anti-HAV titer ≥assay cu	t-off: 20 r	mIU or <del>(</del> 33		
mIU/mL(Enzymun-Test®).				
t anti LIDa A a titar >10 mil	1/201 / 4 1	ICAD®\		

<sup>&</sup>lt;sup>T</sup> anti-HBsAg titer ≥10 mIU/mL (AUSAB<sup>®</sup>).

Since the *Twinrix*-induced Immune responses to hepatitis A and hepatitis B were non-inferior to the monovalent vaccines, efficacy is expected to be similar to the efficacy for each of the monovalent vaccines (Table 3).

Table 3. Geometric Mean Titers in the Twinrix U.S. Clinical Trial

Vaccine	N	Time-point	GMT to Hep A (95% CI)	GMT to Hep B (95% CI)
Twinrix	263	Month 1	335	8
	259	Month 2	636	23
	264	Month 7	4,756 (4,152-5,448)	2,099 (1,663-2,649)
Havrix and Engerix-B	268	Month 1	444	6
	269	Month 2	257	18
	269	Month 7	2,948 (2,638-3,294)	1,871 (1,428-2,450)

It was noted that the antibody titers achieved one month after the final dose of *Twinrix* were higher than titers achieved one month after the final dose of *Havrix* in these clinical trials. This may have been due to a difference in the recommended dosage regimens for these two vaccines, whereby *Twinrix* vaccinees received three doses of 720 EL.U. of hepatitis A antigen at 0, 1 and 6

months, whereas *Havrix* vaccinees received two doses of 1440 EL.U. of the same antigen (at 0 and 6 months). However, these differences in peak titer have not been shown to be clinically significant.

Two clinical trials involving a total of 129 subjects demonstrated that antibodies to both HAV and HBV persisted for at least 4 years after the first vaccine dose in a three-dose series of *Twinrix*, given on a 0-, 1- and 6-month schedule. For comparison, after the recommended immunization regimens for *Havrix* and *Engerix-B*, respectively, similar studies involving a total of 114 subjects have shown that seropositivity to HAV and HBV also persists for at least 4 years.

The effect of age on immune response to *Twinrix* was studied in two trials comparing subjects over 40 years of age (n=183, mean age=48 in one trial and n=72, mean age=50 in the other) with those 40 (n=191; mean age 32.5). The response to the hepatitis A component of *Twinrix* declined slightly with age, but >99% of subjects achieved protective antibody levels in both age groups, and antibody titers were comparable to two doses of hepatitis A vaccine alone in age matched controls.

The response to hepatitis B immunization is known to decline in vaccinees over 40 years of age. Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] elicited a seroprotective response to hepatitis B in 97% of younger subjects and 93% to 94% of the older subjects, as compared to 92% of older subjects given hepatitis B vaccine alone. Geometric mean titers elicited by *Twinrix* were 2,285 in the younger subjects and 1,890 or 1,038 for the older subjects in the two trials. Hepatitis B vaccine alone gave titers of 2,896 in younger subjects and 1,157 in

those over forty.	
It has been shown in open randomized clinical trials that combining the hepatitis A antigen with the hepatitis B surface antigen in <i>Twinrix</i> resulted in comparable anti-HAV or anti-HBsAg titers, relative to vaccination with the individual monovalent vaccines or the concomitant administration of each vaccine in opposite arms.	
Immune response to simultaneously administered vaccines: There have been no studies of concomitant administration of Twinrix with other vaccines.	
INDICATIONS AND USAGE  Twinrix is indicated for active immunization of persons 18 years of age or older against disease caused by hepatitis A virus and infection by all known subtypes of hepatitis B virus. As with any vaccine, vaccination with Twinrix may not protect 100% of recipients. As hepatitis D (caused by the delta virus) does not occur in the absence of HBV infection, it can be expected that hepatitis D will also be prevented by vaccination with Twinrix.	
Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] will not prevent hepatitis caused by other agents such as hepatitis C virus, hepatitis E virus or other pathogens known to infect the liver.	
Immunization is recommended for all susceptible persons 18 years of age or older who are, or will be, at risk of exposure to both hepatitis A and B viruses, including but not limited to:  Travelers to areas of high/intermediate endemicity for both HAV and HBV (see Table 4) who are at increased risk of HBV infection due to behavioral or occupational factors. (See CLINICAL PHARMACOLOGY.)	

Table 4. Hepatitis A and B Endemicity by Region

Geographic Region	HAV	HBV
Africa	High	High (most)
Caribbean	High	Intermediate
Central America	High	Intermediate
South America (temperate)	High	Intermediate
South America (tropical)	High	High
South and Southeast Asia*	High	High
Middle East <sup>†</sup>	High	High
Eastern Europe	Intermediate	Intermediate
Southern Europe	Intermediate	Intermediate
Former Soviet Union	Intermediate	Intermediate

\*Japan: Low HAV and intermediate HBV endemicity † Israel: Intermediate HBV endemicity

Patients with chronic liver disease, including: alcoholic cirrhosis chronic hepatitis C autoimmune hepatitis primary biliary cirrhosis	
Persons at risk through their work: Laboratory workers who handle live hepatitis A and hepatitis B virus Police and other personnel who render first-aid or medical assistance Workers who come in contact with feces or sewage	
Healthcare personnel who render firstaid or emergency medical assistance.	
Personnel employed in day-care centers and correctional facilities. Residents of drug and alcohol treatment centers. Patients and staff of hemodialysis units.	
People living in, or relocating to, areas of high/intermediate endemicity of HAV and who have risk factors for HBV.	
Men who have sex with men.	
Persons at increased risk of disease due to their sexual practices. <sup>20, 21</sup>	
Patients frequently receiving blood products including persons who have clotting-factor disorders (hemophiliacs and other recipients of therapeutic blood products).	

Military recruits and other military	
personnel at increased risk for HBV.	
Users of injectable illicit drugs.	
Individuals who are at increased risk for	
HBV infection and who are close	
household contacts of patients with acute	
or relapsing hepatitis A and individuals	
who are at increased risk for HAV	
infection and who are close household	
contacts of individuals with acute or	
chronic hepatitis B infection.	
CONTRAINIDIOATIONIC	
CONTRAINDICATIONS	
Twinrix [Hepatitis A Inactivated &	
Hepatitis B (Recombinant) Vaccine] is	
contraindicated in people with known hypersensitivity to any component of the	
vaccine or in subjects having shown	
signs of hypersensitivity after previous	
administration of <i>Twinrix</i> or monovalent	
hepatitis A or hepatitis B vaccines.	
Tropanile / Co Tropanile 2 Vaccinics	
WARNINGS	
There have been rare reports of	
anaphylaxis/anaphylactoid reactions	
following routine clinical use of <i>Havrix</i> .	
(See CONTRAINDICATIONS.)	
Hopotitic A and P have relatively long	
Hepatitis A and B have relatively long incubation periods. The vaccine may not	
prevent hepatitis A or B infection in	
individuals who have an unrecognized	
hepatitis A or B infection at the time of	
vaccination. Additionally, it may not	
prevent infection in individuals who do	
not achieve protective antibody titers.	
PRECAUTIONS	
General	
As with other vaccines, although a	
moderate or severe acute illness is	
sufficient reason to postpone	
vaccination, minor illnesses such as mild	
upper respiratory infections with or	
without low-grade fever are not contraindications. <sup>22</sup>	
CONTRAINCICATIONS.	

<b>Pediatric Use:</b> Safety and effectiveness in pediatric patients below the age of 18 years have not been established.	
Geriatric Use: Clinical studies of Twinrix did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects.	
Multiple Sclerosis: Results from two clinical studies indicate that there is no association between hepatitis B vaccination and the development of multiple sclerosis <sup>23</sup> and that vaccination with hepatitis B vaccine does not appear to increase the short-term risk of relapse in multiple sclerosis. <sup>24</sup>	
Twinrix should be administered with caution to people on anticoagulants, those with thrombocytopenia or a bleeding disorder since bleeding may occur following intramuscular administration to these subjects.	
As with any vaccine, if administered to immunosuppressed persons or persons receiving immunosuppressive therapy, the expected immune response may not be obtained. <sup>25</sup>	
Before the injection of any vaccine, the physician should take all reasonable precautions to prevent allergic or other adverse reactions, including understanding the use of the vaccine concerned, and the nature of the side effects and adverse reactions that may follow its use.	
Prior to immunization with any vaccine, the patient's history should be reviewed. The physician should review the patient's immunization history for possible vaccine sensitivity, previous vaccination-related adverse reactions and occurrence of any adverse-event-related symptoms and/or signs, in order to determine the existence	

with T Hepar to allo risks. epine appro imme imme	vincontraindication to immunization winrix [Hepatitis A Inactivated & titis B (Recombinant) Vaccine] and ow an assessment of benefits and As with any parenteral vaccine, phrine injection (1:1000) and other priate agents used for the control of diate allergic reactions must be diately available should an acute hylactic reaction occur.	
a ster each of infe perso	erarate sterile syringe and needle or ille disposable unit must be used for patient to prevent the transmission ectious agents from person to in. Needles should be disposed of erly and should not be recapped.	
Inforr	mation for Patients	
benefit Twinr comp with a subject series concerns advert Department of any to, the Nation 1986.	nts should be informed of the rits and risks of immunization with rix, and of the importance of leting the immunization series. As any vaccine, it is important when a ct returns for the next dose in a sthat he/she be questioned erning the occurrence of any toms and/or signs after a previous of the same vaccine and that se events be reported. The U.S. rtment of Health and Human res has established the Vaccine rese Events Reporting System RS) to accept reports of suspected se events after the administration of vaccine including, but not limited the reporting of events required by the nal Childhood Vaccine Injury Act of The toll-free number for VAERS and information is 1-800-822-	
Pregr	nancy: Pregnancy Category C.	
been not kr fetal h pregn	al reproduction studies have not conducted with <i>T winrix</i> . It is also nown whether <i>Twinrix</i> can cause narm when administered to a nant woman or can affect duction capacity. <i>Twinrix</i> should be	

given to a pregnant woman only if clearly indicated (see INDICATIONS AND USAGE).	
Pregnancy Exposure Registry: Health care providers are encouraged to register pregnant women who receive <i>Twinrix</i> in the SmithKline Beecham Pharmaceuticals vaccination pregnancy registry by calling 1-800-366-8900, ext. 5231.	
Nursing Mothers It is not known whether <i>T winrix</i> is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] is administered to a nursing woman.	
Carcinogenesis, Mutagenesis, Impairment of Fertility  Twinrix has not been evaluated for its carcinogenic potential, mutagenic potential or potential for impairment of fertility.	
ADVERSE REACTIONS In clinical trials involving the administration of 6,594 doses to 2,165 individuals and during routine clinical use of the vaccine outside the United States, <i>Twinrix</i> has been generally well tolerated.	
Of 773 volunteers who participated in the comparative trial conducted in the United States, 389 subjects received at least one dose of <i>Twinrix</i> and 384 received at least one dose each of <i>Engerix-B</i> and <i>Havrix</i> as separate but simultaneous injections. Solicited adverse events reported following the administration of <i>Twinrix</i> are shown in Table 5, compared with adverse events reported after administration of <i>Engerix-B</i> and <i>Havrix</i> .	

Table 5. Rate of Adverse Events Reported After Administration of *Twinrix or Engerix-B* and *Havrix* 

Adverse Event		Twinrix			Engerix-B		На	vrix
Auverse Event	Dose 1	Dose 2	Dose 3	Dose 1	Dose 2	Dose 3	Dose 1	Dose 2
LOCAL	(N=385)	(N=382)	(N=374)	(N=382)	(N=376)	(N=369)	(N=382)	(N=369)
	%	%	%	%	%	%	%	%
Soreness	37	35	41	41	25	30	53	47
Redness	8	9	11	6	7	9	7	9
Swelling	4	4	6	3	5	5	5	5

Adverse Event		Twinrix			Engerix-Band Havrix	
Auverse Everit	Dose 1	Dose 2	Dose 3	Dose 1	Dose 2	Dose 3
GENERAL	(N=385)	(N=382)	(N=374)	(N=382)	(N=376)	(N=369)
	%	%	%	%	%	%
Headache	22	15	13	19	12	14
Fatigue	14	13	11	14	9	10
Diarrhea	5	4	6	5	3	3
Nausea	4	3	2	7	3	5
Fever	4	3	2	4	2	4
Vomiting	1	1	0	1	1	1

Adverse reactions seen with <i>Twinrix</i> were similar to those observed after vaccination with the monovalent components. The frequency of solicited adverse events did not increase with successive doses of Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine]. Most events reported were considered by the subjects as mild and self-limiting and did not last more than 48 hours.	
Among 2,165 subjects in fourteen clinical trials, the following adverse experiences were reported to occur within 30 days following vaccination with the frequency shown below.	
Incidence 1% to 10% of injections  Local reactions at injection site: Induration.  Respiratory system: Upper respiratory tract infections.	

1 11 40/ 61 14	
Incidence <1% of injections	
Local reactions at injection site: Pruritus,	
ecchymoses.	
Body as a whole: Sweating, weakness,	
flushing, influenza-like symptoms.	
Cardiovascular system: Syncope.	
Gastrointestinal system: Abdominal pain,	
anorexia, vomiting.	
Musculoskeletal system: Arthralgia,	
myalgia, back pain.	
Nervous system: Migraine, paresthesia,	
vertigo, somnolence, insomnia, irritability,	
agitation, dizziness.	
Respiratory system: Respiratory tract	
illnesses.	
Skin and appendages: Rash, urticaria,	
petechiae, erythema.	
As with any vaccine, it is possible that	
expanded routine clinical use of the	
vaccine could reveal rare adverse	
events.	
Detential advance offerto	
Potential adverse effects	
The following additional adverse effects	
have been reported with either <i>Engerix-B</i>	
or <i>Havrix</i> in clinical trials and/or during marketed use.	
marketed use.	
Incidence <1% of injections, seen in	
clinical trials with monovalent	
hepatitis A and/or hepatitis B	
vaccines	
Body as a whole: Tingling.b	
Cardiovascular system: Hypotension.b	
Gastrointestinal: Constipation, <sup>b</sup>	
dysgeusia. <sup>a</sup>	
Hematologic/lymphatic:	
Lymphadenopathy. a+b	
Musculoskeletal system: Elevation of	
creatine phosphokinase.a	
Nervous system: Hypertonic episode, <sup>a</sup>	
photophobia. <sup>a</sup>	

Post-marketing Reports	
Since market introduction, more than 61	
million doses of <i>Havrix</i> and more than	
600 million doses of <i>Engerix-B</i> have	
been distributed worldwide (circa	
2000). <sup>27</sup> Voluntary reports of adverse	
events in people receiving either	
Engerix-B or Havrix that have been	
reported since market introduction of the	
vaccines include the following:	
Body as a whole:	
Anaphylaxis/anaphylactoid reactions and	
allergic reactions.a	
Hypersensitivity: Erythema multiforme	
including Stevens-Johnson syndrome, <sup>b</sup>	
angioedema, <sup>b</sup> arthritis, <sup>b</sup> serum-sickness-	
like syndrome days to weeks after	
vaccination including arthralgia/arthritis	
(usually transient), fever, urticaria,	
erythema multiforme, ecchymoses and	
erythema nodosum.b	
Cardiovascular system:	
Tachycardia/palpitations.b	
Skin and appendages: Erythema	
multiforme, a hyperhydrosis, a	
angioedema, <sup>a</sup> eczema, <sup>b</sup> herpes zoster, <sup>b</sup>	
erythema nodosum, <sup>b</sup> alopecia. <sup>b</sup>	
Gastrointestinal system: Jaundice, <sup>a</sup>	
hepatitis, a abnormal liver function tests, b	
dyspepsia. <sup>b</sup>	
Hematologic/ lymphatic:	
Thrombocytopenia. <sup>b</sup>	
Nervous system: Convulsions, a paresis, b	
encephalopathy, a neuropathy, a+b	
myelitis, <sup>a</sup> Guillain-Barré syndrome, <sup>a+b</sup>	
multiple sclerosis, a+b Bell's palsy,b	
transverse myelitis, <sup>b</sup> optic neuritis. <sup>b</sup>	
Respiratory system: Dyspnea, <sup>a</sup>	
bronchospasm including asthma-like	
symptoms.b	
Special senses: Conjunctivitis, b keratitis, b	
visual disturbances, b tinnitus, b earache.b	
Other: Congenital abnormality.a	
a: Following Havrix.	
b: Following Engerix-B.	
a+b: Following either Havrix or Engerix-	
В.	

DOSAGE AND ADMINISTRATION	1
Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] should be administered by intramuscular injection. <i>Do not inject intravenously or intradermally</i> . In adults, the injection should be given in the deltoid region. <i>Twinrix</i> should not be administered in the gluteal region; such injections may result in a suboptimal response.	
For individuals with clotting-factor disorders who are at risk of hemorrhage following intramuscular injection, the ACIP recommends that when any intramuscular vaccine is indicated for such patients, "it should be administered intramuscularly if, in the opinion of a physician familiar with the patient's bleeding risk, the vaccine can be administered with reasonable safety by this route. If the patient receives antihemophilia or other similar therapy, intramuscular vaccination can be scheduled shortly after such therapy is administered. A fine needle (23 gauge or smaller) can be used for the vaccination and firm pressure applied to the site (without rubbing) for at least 2 minutes. The patient should be instructed concerning the risk of hematoma from the injection." <sup>28</sup>	
When concomitant administration of other vaccines or immunoglobulin (IG) is required, they should be given with different syringes and at different injection sites.	
Preparation for Administration: Shake vial or syringe well before withdrawal and use. Parenteral drug products should be inspected visually for particulate matter or discoloration prior to administration. With thorough agitation, <i>Twinrix</i> is a homogenous white turbid suspension. Discard if it appears otherwise.	

The vaccine should be used as supplied; no dilution or reconstitution is necessary. The full recommended dose of the vaccine should be used. After removal of the appropriate volume from a singledose vial, any vaccine remaining in the vial should be discarded.	
Primary immunization for adults consists of three doses, given on a 0-, 1- and 6-month schedule. Each 1 mL dose contains 720 EL.U. of inactivated hepatitis A virus and 20 mcg of hepatitis B surface antigen.	
STORAGE Store refrigerated between 2° and 8° C (36° and 46° F). DO NOT FREEZE; discard if product has been frozen. Do not dilute to administer.	
HOW SUPPLIED Twinrix [Hepatitis A Inactivated & Hepatitis B (Recombinant) Vaccine] is supplied as a turbid white suspension in vials and prefilled Tip-Lok® syringes containing a 1.0 mL single dose.	
Single-Dose Vials NDC 58160-850-01 (package of 1)	
NDC 58160-850-01 (package of 1)	
Single-Dose Prefilled Disposable Tip- Lok® Syringes with 1-inch 23-gauge needles	
NDC 58160-850-35 (package of 5)	
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