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#### HEMOFIL<sup>®</sup> M

#### Antihemophilic Factor (Human), Method M, Monoclonal Purified Nanofiltered

## DESCRIPTION

HEMOFIL M, Antihemophilic Factor (Human) (AHF), Method M, Monoclonal Purified, is a sterile, nonpyrogenic, dried preparation of antihemophilic factor (Factor VIII, Factor VIII:C, AHF) in concentrated form with a specific activity range of 2 to 22 AHF International Units/mg of total protein. HEMOFIL M contains a maximum of 12.5 mg/mL Albumin, and per AHF International Unit, 0.07 mg polyethylene glycol (3350), 0.39 mg histidine as stabilizing agents, not more than 0.1 mg glycine, 0.1 ng mouse protein, 18 ng organic solvent (tri-n-butyl phosphate) and 50 ng detergent (octoxynol 9). In the absence of the added Albumin (Human), the specific activity is approximately 2,000 AHF International Units/mg of protein [see Clinical Pharmacology].

HEMOFIL M is prepared by the Method M process from pooled human plasma by immunoaffinity chromatography utilizing a murine monoclonal antibody to Factor VIII:C, followed by an ion exchange chromatography step for further purification. Source material may be provided by other US licensed manufacturers. HEMOFIL M also includes an organic solvent (tri-n-butyl phosphate) and detergent (octoxynol 9) virus inactivation step designed to reduce the risk of transmission of hepatitis and other viral diseases. The process further includes a nanofiltration step between immunoaffinity chromatography and ion-exchange chromatography as an additional viral clearance step to further improve the viral safety margin of the final product.

Use of an organic solvent (tri-n-butyl phosphate; TNBP) in the manufacture of Antihemophilic Factor (Human) has little or no effect on AHF activity, while lipid enveloped viruses, such as hepatitis B and human immunodeficiency virus (HIV) would be inactivated.<sup>1</sup> The nanofiltration step integrated into the manufacture of AHF-M further enhances the safety margin with respect to adventitious viruses. Each bottle of HEMOFIL M is labeled with the AHF activity expressed in International Units (IU) per bottle. This potency assignment is referenced to the World Health Organization International Standard. The purity of HEMOFIL M has been thought to influence the difficulty of producing an accurate potency measurement. Experiments have shown that to achieve accurate activity levels, such a potency assay should be conducted using plastic test tubes and pipets as well as substrate containing normal levels of von Willebrand's Factor.

*In vitro* studies demonstrate that the HEMOFIL M manufacturing process provides for viral reduction. These reductions are achieved through a combination of process chemistry, partitioning and/or inactivation during solvent/detergent treatment, and immunoaffinity chromatography. Introduction of a nanofiltration step with the 0.1µm prefilter and the ASAHI Planova 20N nanofilter provides a virus removal capacity for human immunodeficiency virus, Type 1 (HIV-1), hepatitis A virus (HAV), bovine viral

diarrhea virus (BVDV), pseudorabies virus (PRV), mice minute virus (MMV), and human parvovirus B19 (B19V) in the order of four (4) logs or higher. B19V removal data were obtained with a Polymerase Chain Reaction (PCR) assay not correlated to an infectivity assay.

Studies for nanofiltration and other process steps, summarized in Table 1, demonstrate virus clearance during the HEMOFIL M manufacturing process using HIV-1; BVDV, a generic model for lipid enveloped RNA viruses, such as hepatitis C virus (HCV); PRV, a model for lipid enveloped DNA viruses, such as hepatitis B virus (HBV); canine parvovirus (CPV), a model for non-lipid enveloped DNA viruses, such as B19V, HAV, and MMV.

Dracase Star	Virus Clearance, log <sub>10</sub>					
Evaluated	Lipid-enveloped			Non-Lipid enveloped		
	HIV-1	BVDV	PRV	HAV	CPV	MMV
Solvent/Detergent Treatment	>4.8	>6.8	>6.9	$NT^*$	NT*	NT*
Immunoaffinity Chromatography	N.A. <sup>†</sup>	N.A.†	N.A.†	≥4.5	≥3.9	NT
Nanofiltration	>5.5	>4.6	>4.4	>5.4	NT	>5.0
Cumulative Total, log <sub>10</sub>	>10.3	>11.4	>11.3	>9.9	≥3.9	>5.0
NT						

Table 1 In Vitro Virus Clearance During the Manufacture of HEMOFIL M

NT not tested

\* As Solvent/Detergent treatment does not inactivate non-lipid enveloped viruses.

+ Not Applicable for lipid enveloped viruses due to the presence of (virucidal) solvent/detergent reagents in the starting material.

# CLINICAL PHARMACOLOGY

Antihemophilic factor (AHF) is a protein found in normal plasma which is necessary for clot formation. The administration of HEMOFIL M provides an increase in plasma levels of AHF and can temporarily correct the coagulation defect of patients with hemophilia A (classical hemophilia). The half-life of HEMOFIL M administered to Factor VIII deficient patients has been shown to be 14.8  $\pm$  3.0 hours.

# INDICATIONS AND USAGE

HEMOFIL M is indicated in hemophilia A (classical hemophilia) for the prevention and control of hemorrhagic episodes.

HEMOFIL M is not indicated in von Willebrand's disease.

# CONTRAINDICATIONS

HEMOFIL M is contraindicated in patients with a known hypersensitivity to the active substance, to excipients, or to mouse proteins.

## WARNINGS

## Hypersensitivity

Allergic-type hypersensitivity reactions, including anaphylaxis, have been reported with HEMOFIL M and have been manifested by bronchospasm, dyspnea, hypotension, chest pain, facial edema, urticaria, rash, flushing, pruritus, and nausea.

### **Neutralizing Antibodies**

The development of neutralizing antibodies (inhibitors) to Factor VIII is a known complication of the treatment of patients with Hemophilia A. Inhibitors have predominantly been reported in previously untreated patients. The risk of developing inhibitors is correlated to the extent of exposure to Factor VIII, the risk being highest within the first 20 exposure days, and to other genetic and environmental factors. The risk for inhibitor development depends on a number of factors relating to the characteristics of the patient (e.g., type of the Factor VIII gene mutation, family history, ethnicity), which are believed to represent the most significant risk factors for inhibitor formation.

## **Transmission of Infectious Agents**

Because HEMOFIL M is made from human blood, it may carry a risk of transmitting infectious agents, e.g., viruses, the variant Creutzfeldt-Jakob disease (vCJD) agent and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent. This also applies to unknown or emerging viruses and other pathogens.

All infections thought by a physician possibly to have been transmitted by this product should be reported by the physician or other healthcare provider to Takeda Pharmaceuticals U.S.A., Inc. at 1-877-TAKEDA-7 (1-877-825-3327) or FDA at 1-800-FDA-1088 or *www.fda.gov/medwatch*. The physician should discuss the risks and benefits of this product with the patient.

Individuals who receive infusions of blood or plasma products may develop signs and/or symptoms of some viral infections, particularly non-A, non-B hepatitis. As indicated under *Clinical Pharmacology*, however, a group of such patients treated with HEMOFIL M did not demonstrate signs or symptoms of non-A, non-B hepatitis over observation periods ranging from three to nine months.

## PRECAUTIONS

Identification of the clotting defect as a Factor VIII deficiency is essential before the administration of HEMOFIL M is initiated.

#### Factor VIII Inhibitors

Evaluate patients for the development of Factor VIII inhibitors if the expected plasma Factor VIII activity levels are not attained or if bleeding is not controlled with an appropriate dose.

No benefit may be expected from this product in treating other deficiencies.

## Formation of Antibodies to Mouse Protein

HEMOFIL M contains trace amounts of mouse protein (less than 0.1 ng/AHF activity units). The possibility exists that patients treated with HEMOFIL M may develop hypersensitivity to the mouse proteins. There have been no cases of hypersensitivity to the mouse proteins reported.

# Increase in Pulse Rate

Determine the pulse rate before and during administration of HEMOFIL M. Should a significant increase occur, reduce the rate of administration or temporarily halt the injection to allow the symptoms to disappear promptly.

# Laboratory Tests

Perform appropriate laboratory tests on the patient's plasma at suitable intervals to ensure that adequate AHF levels have been reached and are maintained.

If the AHF content of the patient's plasma fails to reach expected levels or if bleeding is not controlled after apparently adequate dosage, the presence of inhibitor should be suspected. By appropriate laboratory procedures, the presence of inhibitor can be demonstrated and quantified in terms of AHF units neutralized by each mL of plasma or by the total estimated plasma volume.

If the inhibitor is at low levels (i.e., <10 Bethesda units per mL), after administration of sufficient AHF units to neutralize the inhibitor, additional AHF units will elicit the predicted response.

# Pregnancy

Animal reproduction studies have not been conducted with HEMOFIL M. The safety of HEMOFIL M for use in pregnant women has not been established. It is not known whether HEMOFIL M can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. HEMOFIL M should be given to a pregnant woman only if clearly needed.

# Nursing Mothers

The safety of HEMOFIL M for use in nursing mothers has not been established. It is not known whether this drug is excreted into human milk. Physicians should carefully consider the potential risks and benefits for each specific patient before prescribing HEMOFIL M. HEMOFIL M should be given to nursing mothers only if clinically indicated.

# ADVERSE REACTIONS

# Adverse Reactions from Clinical Trials

The adverse reactions presented in this section have been identified based on clinical trial experience with HEMOFIL M in patients previously treated with other Factor VIII concentrates or blood products (N = 74), and previously untreated patients (PUPs; N = 50).

		(Frequency Percentage)
BLOOD AND LYMPHATIC SYSTEM DISORDERS	Factor VIII inhibition	3 (5.7%)*
NERVOUS SYSTEM	Dizziness	1 (0.8%)
DISORDERS	Headache	1 (0.8%)
	Dysgeusia	1 (0.8%)
GENERAL DISORDERS AND	Pyrexia	1 (0.8%)
ADMINISTRATION SITE	Infusion site inflammation	2 (1.6%)

\* In a study that included 43 evaluable PUPs and 10 minimally treated patients (MTPs), i.e., patients with a single exposure to other Factor VIII concentrates or blood products, 3 of the total of 53 patients (5.7%) developed an inhibitor while on study.

HEMOFIL M was administered to 11 patients previously untreated with Antihemophilic Factor (Human). They have shown no signs of hepatitis or HIV infection following three to nine months of evaluation.

A study of 25 patients treated with HEMOFIL M and monitored for three to six months has demonstrated no evidence of antibody response to mouse protein. More than 1,000 infusions of HEMOFIL M have been administered during the clinical trials. Reported events included a single episode each of chest tightness, fuzziness and dizziness, and one patient reported an unusual taste after each infusion.

## **Post-marketing Adverse Reactions**

In addition to clinical trials, the following adverse reactions have been reported in the post-marketing experience, listed by MedDRA System Organ Class (SOC), then by Preferred Term.

Immune System Disorders: anaphylactic reaction, hypersensitivity

Eye Disorders: visual impairment, ocular hyperemia

Cardiac Disorders: cyanosis, bradycardia, tachycardia

Vascular Disorders: hypotension, flushing

<u>Respiratory, Thoracic, and Mediastinal Disorders</u>: bronchospasm, dyspnea, cough, hyperventilation

Gastrointestinal Disorders: diarrhea, vomiting, nausea, abdominal pain

Skin and Subcutaneous Tissue Disorders: urticaria, rash, pruritus, hyperhidrosis

Musculoskeletal and Connective Tissue Disorders: musculoskeletal pain

<u>General Disorders and Administration Site Conditions</u>: facial edema, edema, chills, fatigue, chest pain, irritability

# DOSAGE AND ADMINISTRATION

For intravenous use only.

The expected in vivo peak AHF level, expressed as IU/dL of plasma or % (percent) of

normal, can be calculated by multiplying the dose administered per kg body weight (IU/kg) by two. This calculation is based on the clinical finding by Abildgaard, *et al*,<sup>2</sup> which is supported by data from the collaborative study of *in vivo* recovery and survival with 15 different lots of HEMOFIL M on 56 hemophiliacs that demonstrated a mean peak recovery point above the mean pre-infusion baseline of about 2.0 IU/dL per infused IU/kg body weight.<sup>3</sup>

Examples:

- (1) A dose of 1750 IU AHF administered to a 70 kg patient, i.e., 25 IU/kg (1750/70), should be expected to cause a peak post-infusion AHF increase of 25 x 2 = 50 IU/dL (50% of normal).
- (2) A peak level of 70% is required in a 40 kg child. In this situation the dose would be  $70/2 \times 40 = 1400 \text{ IU}$ .

# **Recommended Dosage Schedule**

Physician supervision of the dosage is required. The following dosage schedule may be used as a guide.

HEMORRHAGE		
Degree of hemorrhage	Required peak post- infusion AHF activity in the blood (as % of normal or IU/dL plasma)	Frequency of infusion
Early hemarthrosis or muscle bleed or oral bleed	20-40	Begin infusion every 12 to 24 hours for one-three days until the bleeding episode as indicated by pain is resolved or healing is achieved.
More extensive hemarthrosis, muscle bleed, or hematoma	30-60	Repeat infusion every 12 to 24 hours for usually three days or more until pain and disability are resolved.
Life threatening bleeds such as head injury, throat bleed, severe abdominal pain	60-100	Repeat infusion every 8 to 24 hours until threat is resolved.
SURGERY	Г <u> </u>	
Type of operation		
Minor surgery, including tooth extraction	60-80	A single infusion plus oral antifibrinolytic therapy within one hour is sufficient in approximately 70% of cases.
Major surgery	80-100 (pre- and post-operative)	Repeat infusion every 8 to 24 hours depending on state of healing.

If bleeding is not controlled with the prescribed dose, determine the plasma level of Factor VIII and administer a sufficient dose of HEMOFIL M to achieve a satisfactory clinical response. Under certain circumstances (e.g., presence of a low titer inhibitor) doses larger than those recommended may be necessary as per standard care. In patients with high titer Factor VIII inhibitors, HEMOFIL M therapy may not be effective and other therapeutic options should be considered. The dosage and duration of treatment depend on the severity of Factor VIII deficiency, the location and extent of the bleeding, and the patient's clinical condition. Careful control of replacement therapy is especially important in cases of major surgery or life threatening hemorrhages.

# **Reconstitution: Use Aseptic Technique**

- 1. Bring HEMOFIL M (dry concentrate) and Sterile Water for Injection, USP, (diluent) to room temperature.
- 2. Remove caps from concentrate and diluent bottles to expose central portion of rubber stoppers.
- 3. Cleanse stoppers with germicidal solution.
- 4. Remove protective covering from one end of double-ended needle and insert exposed needle through diluent stopper.
- 5. Remove protective covering from other end of double-ended needle. Invert diluent bottle over upright HEMOFIL M bottle, then rapidly insert free end of the needle through the HEMOFIL M bottle stopper at its center. The vacuum in the HEMOFIL M bottle will draw in the diluent.
- 6. Disconnect the two bottles by removing needle from diluent bottle stopper, then remove needle from HEMOFIL M bottle. Swirl gently until all material is dissolved. Be sure that HEMOFIL M is completely dissolved; otherwise active material will be removed by the filter.

Note: Do not refrigerate after reconstitution.

# Administration: Use Aseptic Technique

- Intravenous administration only.
- Administer at room temperature not more than 3 hours after reconstitution.
- Record the name and batch number of the product in order to maintain a link between the patient and the batch of the product.

# Intravenous Syringe Injection

- Visually inspect parenteral product for particulate matter and discoloration prior to administration. The solution should be colorless in appearance. Do not administer if particulate matter or discoloration is found.
- Plastic syringes are recommended for use with this product. The ground glass surface of all-glass syringes tend to stick with solutions of this type.
- 1. Attach filter needle to a disposable syringe and draw back plunger to admit air into syringe.
- 2. Insert needle into reconstituted HEMOFIL M.
- 3. Inject air into bottle and then withdraw the reconstituted material into the syringe.
- 4. Remove and discard the filter needle from the syringe; attach a suitable needle and inject intravenously as instructed under *Rate of Administration*.
- 5. If a patient is to receive more than one bottle of HEMOFIL M, the contents of two bottles may be drawn into the same syringe by drawing up each bottle through a

separate unused filter needle. This practice lessens the loss of HEMOFIL M. Filter needles are intended to filter the contents of a single bottle of HEMOFIL M only.

## **Rate of Administration**

Administer HEMOFIL M at a rate of up to 10 mL per minute. Infuse HEMOFIL M at a rate of administration that ensures the comfort of the patient [see Precautions: Increase of Pulse Rate].

## HOW SUPPLIED

HEMOFIL M is available as single-dose bottles that contain the following nominal potencies:

Nominal Potency	Kit NDC Number
250 IU	0944-3940-02
500 IU	0944-3942-02
1000 IU	0944-3944-02
1700 IU	0944-3946-02

Each bottle is labeled with the potency in International Units, and is packaged together with 10 mL of Sterile Water for Injection, USP, a double-ended needle, and a filter needle.

Not made with natural rubber latex.

# Storage

HEMOFIL M can be stored at 2°C to 8°C (36°F to 46°F) or at room temperature, not to exceed 30°C (86°F), until expiration date noted on the package.

Do not freeze.

## Information for Patients

- Advise patients to report any adverse reactions or problems following HEMOFIL M administration to their physician or healthcare provider.
- Advise pregnant women or immune compromised individuals of the effects of Parvovirus B19. Symptoms include fever, drowsiness, chills, runny nose followed about two weeks later by a rash, and joint pain.
- Inform patients of the signs and symptoms of hepatitis A, which include several days to weeks of poor appetite, tiredness, and low-grade fever followed by nausea, vomiting, and pain in the belly. Dark urine and a yellowed complexion are also common symptoms. Encourage patients to consult their physician if such symptoms appear.
- Inform patients of the early signs of hypersensitivity reactions including hives, generalized urticaria, facial edema, flushing, nausea, tightness of the chest, wheezing, dyspnea, hypotension, and anaphylaxis. Advise patients to discontinue use of the product and contact their physician if these symptoms occur.

## REFERENCES

- 1. Horowitz B, Wiebe ME, Lippin A, *et al*: Inactivation of viruses in labile blood derivatives: 1. Disruption of lipid enveloped viruses by tri(n-butyl)phosphate detergent combinations. **Transfusion 25**:516-522, 1985.
- 2. Abildgaard CF, Simone JV, Corrigan JJ, *et al*: Treatment of hemophilia with glycineprecipitated Factor VIII. **New Eng J Med 275**:471-475, 1966.
- 3. Addiego, Jr. JE, Gomperts E, Liu S. *et al*: Treatment of hemophilia A with a highly purified Factor VIII concentrate prepared by Anti-FVIIIc immunoaffinity chromatography. **Thrombosis and Haemostasis 67**:19-27, 1992.

To enroll in the confidential, industry-wide Patient Notification System, call 1-888-873-2838.

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Revised: 2/2025

## PRINCIPAL DISPLAY PANEL - 10 mL Bottle Label - 250 iU

10 mL size, dried List 1502845 NDC 0944-3941-01

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

## FVIII

## Intravenous administration only

Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C (86°F) until expiration date noted on package.

Dosage and Administration: read package insert. Contains no preservative.

# **Rx Only**

Takeda Pharmaceuticals U.S.A., Inc. Cambridge, MA 02142 U.S. License No. 1898

23		
Antihemophilic Factor (Human), Method M. Monoclonal Purified	20.mL size, dried List 1502845 NDC 0944-3941-01	
HEMOFIL* M Nanofiltered NDC 0944-3941-01	Antihemophilic Factor (Human)	
Date of Dose:		
vial:/	8 Nanofiltered	
	Intravenous administration only	
LOT NO.:	→ Store at 2° - 8°C (36° - 46°F) or room temperature, ⊂ not to exceed 30°C (86°F) until expiration date	
Time of Dose:	C noted on package.	
Takeda Pharmaceuticals U.S.A., Inc.	Contains no preservative.	
Cambridge, MA 02142	Takeda Pharmaceuticals U.S.A., Inc.	
Peel At Arrow for Patient Records 6521204		

# PRINCIPAL DISPLAY PANEL - Kit Carton - 250 iU

10 mL size, dried

NDC 0944-3940-02

#### Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

## FVIII

## Intravenous administration only.

Administer within 3 hours after reconstitution.

Dosage and Administration: Read full prescribing information.

Warning: This product is prepared from large pools of human plasma. Human blood and its components may transmit infectious agents. The patient and the physician should discuss the risks and benefits of this product.

Contains no preservative.

**Rx Only** 

10 mL size, dried	NDC 0944-3940-02	
Antihemophilic Factor (Human)		Antihemophilic Factor (Human)
Method M, Monoclonal Purified		Method M, Monoclonal Purified HEMOFIL® M
HEMOFIL® M Napofiltered	FVIII	Nanofiltered FVIII
Natonicieu		Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C
Intravenous administration only.		(86°F) until expiration date noted on the package.
Administer within 3 hours after reconstitution.		Do not freeze.
Dosage and Administration: Read full prescribing information.		Date removed from refrigeration and
Warning: This product is prepared from large pools of human plasma	a. Human blood and	placed at room temperature.
its components may transmit infectious agents. The patient should discuss the risks and benefits of this product.	t and the physician	Date: / /
Contains no preservative.	-	Date///
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Product Cod et 1502845	Antihern op hilic Factor (Human) Method M, Monoclonal Purified Manofilered Eviliuv Exp. Date: Exp. Date:	
Product Code: 1502845 Antihemophilic Factor (Hur Method M, Monoclonal Pur HEMOFIL® M Nanofiltered Contents: One 10 mL bottle dried Antihe Water for Injection, USP; one double-en- insert. Stabilizing agents present in the followin (Human) and per AHF International Unit contains not more than 0.1 mg glycine, (tri-n-butyl phosphate) and 50 ng deterg HEMOFIL M, double-ended needle and filte Sterile Water product of Germany. Takeda Pharmaceuticals U.S.A., Inc. Cambridge, MA 02142 U.S. License No. 1898	nan) rified FOUL mophilic Factor (Human); 10 mL Sterile ded needle; one filter needle; and package mg maximum amounts: 12.5 mg/mL Albumin , 0.07 mg PEG, 0.39 mg histidine, and it also 0.1 ng mouse protein, 18 ng organic solvent gent (octoxynol 9). r needle products of USA;	Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered FVIII Sterile Nonpyrogenic Sterile Nonpyrogenic Components are not made with natural rubber latex. To enroll in the confidential, industry-wide Patient Notification System, call 1-888-873-2838. HEMOFIL is a registered trademark of Baxaita Incorporated. Takeda and are registered trademarks of Takeda Pharmaceutical Company Limited.

# PRINCIPAL DISPLAY PANEL - 10 mL Bottle Label - 500 iU

10 mL size, dried List 1502846 NDC 0944-3943-01

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered

## FVIII

## Intravenous administration only

Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C (86°F) until expiration date noted on package.

Dosage and Administration: read package insert. Contains no preservative.

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# Takeda



# PRINCIPAL DISPLAY PANEL - Kit Carton - 500 iU

10 mL size, dried

NDC 0944-3942-02

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

# FVIII

## Intravenous administration only.

Administer within 3 hours after reconstitution.

Dosage and Administration: Read full prescribing information.

Warning: This product is prepared from large pools of human plasma. Human blood and its components may transmit infectious agents. The patient and the physician should discuss the risks and benefits of this product.

Contains no preservative.

**Rx Only** 

10 mL size, dried Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered Intravenous administration only. Administer within 3 hours after reconstitution. Dosage and Administration: Read full prescribing information. Warning: This product is prepared from large pools of human plasm its components may transmit infectious agents. The patient should discuss the risks and benefits of this product.	NDC 0944-3942-02 FVIII na. Human blood and nt and the physician	Antihemophilic Factor (Human)   Method M, Monoclonal Purified   HEMOFIL® M   Nanofiltered   Store at 2° - 8°C (36° - 46°F) or room   temperature, not to exceed 30°C   (86°F) until expiration date noted on   the package.   Do not freeze.   Date removed from refrigeration and   placed at room temperature.
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Product Cod & 1502846	Antihemophilic Factor (Human) Method M, Monoclonal Purified M ® JFA Monoclonal Purified Manofiltered Evile: Lot Vo.: Exp. Date:	
Product Code: 1502846 Antihemophilic Factor (Hum Method M, Monoclonal Purit HEMOFIL® M Nanofiltered Contents: One 10 mL bottle dried Antihem Water for Injection, USP; one double-ender insert. Stabilizing agents present in the following (Human) and per AHF International Unit, O contains not more than 0.1 mg glycine, 0. (tri-n-butyl phosphate) and 50 ng deterge	an) fied FVIII pophilic Factor (Human); 10 mL Sterile ad needle; one filter needle; and package maximum amounts: 12.5 mg/mL Albumin 0.07 mg PEG, 0.39 mg histidine, and it also 1 ng mouse protein, 18 ng organic solvent nt (octoxynol 9).	Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered FVIII Sterile Nonpyrogenic Components are not made with natural rubber latex. To enroll in the confidential, industry-wide Patient Notification System, call 1-888-873-2838. HEMOFIL is a registered trademark of Baxalta Incorporated. Takeda and Amagina are registered

## PRINCIPAL DISPLAY PANEL - 10 mL Bottle Label - 1000 iU

10 mL size, dried List 1502847 NDC 0944-3945-01

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

# FVIII

### Intravenous administration only

Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C (86°F) until expiration date noted on package.

Dosage and Administration: read package insert. Contains no preservative.

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## Takeda



# PRINCIPAL DISPLAY PANEL - Kit Carton - 1000 iU

10 mL size, dried

NDC 0944-3944-02

#### Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

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## FVIII

# Intravenous administration only.

Administer within 3 hours after reconstitution.

Dosage and Administration: Read full prescribing information.

Warning: This product is prepared from large pools of human plasma. Human blood and

its components may transmit infectious agents. The patient and the physician should discuss the risks and benefits of this product.

Contains no preservative.

**Rx Only** 

10 mL size, dried	NDC 0944-3944-02	
Antihemophilic Factor (Human) Method M, Monoclonal Purified		Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M
HEMOFIL® M	FVIII	Nanofiltered FVIII
Nanomered		Store at 2° - 8°C (36° - 46°F) or room
Intravenous administration only.		temperature, not to exceed 30°C (86°F) until expiration date noted on
Administer within 3 hours after reconstitution.		the package.
Dosage and Administration: Read full prescribing information.		Do not freeze.
Warning: This product is prepared from large pools of human plas its components may transmit infectious agents. The pati should discuss the risks and benefits of this product.	sma. Human blood and ent and the physician	Date removed from refrigeration and placed at room temperature.
Contains no preservative.	-	Date://
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Product Code: 1502847	Antih emophilic Factor (Human) Method M, Monoclonal Puritied FVIII IU / Bottle: Lot No.: Exp. Date: Exp. Date:	
Product Code: 1502847 Antihemophilic Factor (Huma Method M, Monoclonal Purifi HEMOFIL® M Nanofiltered Contents: One 10 mL bottle dried Antihemo Water for Injection, USP; one double-ender insert. Stabilizing agents present in the following (Human) and per AHF International Unit, 0 contains not more than 0.1 mg glycine, 0.1 (tri-n-butyl phosphate) and 50 ng detergen HEMOFIL M, double-ended needle and filter n Sterile Water product of Germany. Takeda Pharmaceuticals U.S.A., Inc. Cambridge, MA 02142 U.S. License No. 1898	nn) ied <b>EVIII</b> pophilic Factor (Human); 10 mL Sterile d needle; one filter needle; and package maximum amounts: 12.5 mg/mL Albumin 07 mg PEG, 0.39 mg histidine, and it also ng mouse protein, 18 ng organic solvent t (octoxynol 9). eedle products of USA;	Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered FULL Sterile Nonpyrogenic Components are not made with natural rubber latex. To enroll in the confidential, industry-wide Patient Notification System, call 1-888-873-2838. HEMOFIL is a registered trademark of Baxalta Incorporated. Takeda and are registered trademarks of Takeda Pharmaceutical Company Limited. @2025 Takeda Pharmaceutical Company Limited. All rights reserved.

## PRINCIPAL DISPLAY PANEL - 10 mL Bottle Label - 1700 iU

10 mL size, dried List 1502848 NDC 0944-3947-01

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

### FVIII

## Intravenous administration only

Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C (86°F) until expiration date noted on package.

Dosage and Administration: read package insert. Contains no preservative.

## **Rx Only**

Takeda Pharmaceuticals U.S.A., Inc. Cambridge, MA 02142 U.S. License No. 1898

## Takeda



# PRINCIPAL DISPLAY PANEL - Kit Carton - 1700 iU

10 mL size, dried

NDC 0944-3946-02

Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL<sup>®</sup> M Nanofiltered

## FVIII

## Intravenous administration only.

Administer within 3 hours after reconstitution.

Dosage and Administration: Read full prescribing information.

Warning: This product is prepared from large pools of human plasma. Human blood and its components may transmit infectious agents. The patient and the physician should discuss the risks and benefits of this product. Contains no preservative.

**Rx Only** 

10 mL size, dried Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered Intravenous administration only. Administer within 3 hours after reconstitution. Dosage and Administration: Read full prescribing information. Warning: This product is prepared from large pools of human points components may transmit infectious agents. The poly should discuss the risks and benefits of this product. R. Only	NDC 0944-3946-02	Antihemophilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M Nanofiltered Store at 2° - 8°C (36° - 46°F) or room temperature, not to exceed 30°C (86°F) until expiration date noted on the package. Do not freeze. Date removed from refrigeration and placed at room temperature.
0 309443 946020		6521279
6251529		

Product Code: 1502848	Antih emop hilic Factor (Human) Method M, Monoclonal Purified HEMOFIL® M FVIII IU / Botte: Lot No.: Exp. Date:	

# PRINCIPAL DISPLAY PANEL - 10 mL Vial Label

10 mL

NDC 64764-516-10

## Takeda

Sterile Water for Injection, USP for reconstitution of accompanying product

Do not use unless clear. No antimicrobial agent or other substance has been added. Do not use for intravascular injection without making approximately isotonic by addition of suitable solute. Discard unused portion. **Rx Only** 

Single-dose container Nonpyrogenic

6521215

Product	Manufact	10 mL	NDC 64764-516-10	Takeda	6109			
of Germany	tured for:	Sterile for recons	Water for Institution of accor	njection, USP	56476451			
2 2 2		Do not use un has been addeo approximately Discard unuse	less clear. No antimicrob d. Do not use for intravas isotonic by addition of d portion. <b>Rx Only</b>	ial agent or other substance cular injection without making suitable solute. Single-dose container				
II.	20		6521215	Nonpyrogenic		LOT	EXP	

HEMOFIL M antihemophilic factor human kit								
Produ	ıct Informa	tion						
Product Type   PLAS MA DERIVATIVE		Item Code (Source)	NDC:0944-3940					
Packa	Packaging							
# It	em Code	<b>Package Description</b>	Marketing Start Date	Marketing End Date				
1 NDC:0	0944-3940-02	1 in 1 CARTON						
1								
Quant	tity of Parts	5						
Part #	Pa	ackage Quantity	Total Product Quantity					
Part 1	1 BOTTLE		10 mL					
Part 2	1 VIAL, GLASS		10 mL					
Part	Part 1 of 2							
<b>HEM</b> antiher	<b>OFIL M</b> mophilic facto	or human powder, for soluti	on					

Product Information							
Item Code (Source	e)	NDC:0944-3941					
Route of Administ	ration	INTRAVENOUS					
Active Ingredier	nt/Active	Moietv					
jj	ength	Strength					
ANTIHEMOPHILIC FA		AN (UNII: 839MOZ74GK) (ANTIHEMOF	PHILIC	ANTIHEMOPHILI	C	250 [iU]	
FACTOR HUMAN - UNII:	839MOZ 74GI	<)		FACTOR HUMAN	J	in 10 mL	
Inactive Ingredi	ents						
		Ingredient Name			St	rength	
ALBUMIN HUMAN (UN	NII: ZIF514RV	ZR)					
	COL 3350 (l	JNII: G2M7P15E5P)					
	59/98/E)						
Packaging							
# Item	Item Backage Description Marketin					larketing	
" Code			instisu	Start Da	te I	End Date	
<b>1</b> NDC:0944- 3941-01 Product	t (e.g., Drug/	Device/Biological Product)	ination				
Marketing In	format	ion					
Marketing Category	Applicat	tion Number or Monograph Citation	Marke	ting Start Date	Marketing End Date		
BLA	BLA101448		02/23/198	38			
Part 2 of 2							
STERILE WAT	ΓER						
water liquid							
<b>Product Inform</b>	ation						
Item Code (Source) NDC:64764-516							
Route of Administ	ration	INTRAVENOUS					
Inactive Ingredi	onto						
mactive ingredi	Ingrad	iont Namo		Stro	nath		
	KOOR)		10 ml	. in 10 ml	iyui		
			10 111	10			

Pa	Packaging								
#	ltem Code		Package Description	Market Start Da	ing ate	Marketing End Date			
1	NDC:64764- 516-10	10 mL Produc	in 1 VIAL, GLASS; Type 9: Other Type of Part 3 t (e.g., Drug/Device/Biological Product)	Combination					
Μ	larketin	ng In	formation						
	Marketin Categor	ng 'Y	Application Number or Monograph Citation	Marketing Start Date		Marketing End Date			
BL	A		BLA101448	02/23/1988					
Marketing Information									
	Marketin Categor	ng 'Y	Application Number or Monograph Citation	Marketin Dat	g Start e	Ма	rketing End Date		
BL	A		BLA101448	02/23/1988					

HEMOFIL M antihemophilic factor human kit							
Product Information							
Product Type	PLASMA DERIVATIVE	ltem Code (Source)	NDC:0944-3942				
Packaging							
# Item Code	Package Description	Marketing Start Date	Marketing End Date				
<b>1</b> NDC:0944-3942-02	1 in 1 CARTON						
<b>Quantity of Parts</b>	5						
Part # Pa	ackage Quantity	Total Produ	ict Quantity				
Part 1 BOTTLE		10 mL					
Part 2 1 VIAL, GLASS		10 mL					
Part 1 of 2							
HEMOFIL M antihemophilic factor human powder, for solution							
Product Informa	tion						

Item Code (	Source	2)	NDC:0944-3943				
Route of Ad	minist	ration	INTRAVENOUS				
Active Ing	redier	nt/Active	Moiety				
		Ingre	edient Name		Basis of St	rength	Strength
ANTIHEMOPH FACTOR HUMAN	ILIC FA N - UNII:	CTOR HUMA 839MOZ 74GI	<b>AN</b> (UNII: 839MOZ74GK) (ANTIHEMOI <)	PHILIC	ANTIHEMOPHII FACTOR HUMA	LIC N	500 [iU] in 10 mL
Inactive In	gredi	ents					
			Ingredient Name			St	rength
ALBUMIN HUN	MAN (UN	NII: ZIF514RVZ	ZR)				
POLYETHYLE	NE GLY	COL 3350 (l	JNII: G2M7P15E5P)				
HISTIDINE (UN	III: 4QD	397987E)					
Packaging							
# Item Code			Package Description		Marketi Start Da	ing M ate I	Aarketing End Date
<b>1</b> NDC:0944- 3943-01	10 mL Product	in 1 BOTTLE; t (e.g., Drug/	Type 9: Other Type of Part 3 Comb Device/Biological Product)	ination			
Marketin	ng In	formati	ion				
Marketi Catego	ng ry	Applicat	tion Number or Monograph Citation	Marke	ting Start Date	Mark	eting End Date
BLA		BLA101448		02/23/198	38		
Part 2 of	f 2						
STERILE	WA	ΓER					
water liquid							
Product In	nform	ation					
Item Code (	Source	e)	NDC:64764-516				
Route of Ad	minist	ration	INTRAVENOUS				
Inactive In	gredi	ents					
	-	Ingredi	ent Name		Stre	ength	
WATER (UNII:	059QF0	KO0R)		10 mL	in 10 mL		

Da	a ka a ina									
ra	CKaying							Marke	• :	
#	Code			Package De	escrip	otion		Start D	ting Date	End Date
1 5	IDC:64764- 16-10	10 mL Produc	in 1 VIAL, GL t (e.g., Drug	ASS; Type 9: Otl /Device/Biologica	her Ty al Prod	pe of Part 3 C uct)	ombination			
Ma	arketir	ng In	format	ion						
	Marketin Categor	ng 'Y	Applicat	tion Number o Citation	or Mo n	nograph	Marketin Dat	g Start e	Ma	rketing End Date
BLA			BLA101448				02/23/1988			
Ma	arketir	ng In	format	ion						
	Marketin Categor	ng 'Y	Applicat	tion Number of Citation	or Mo n	nograph	Marketin Dat	g Start e	Ma	rketing End Date
BLA			BLA101448				02/23/1988			
HE	HEMOFIL M									
antil	antihemophilic factor human kit									
Pro	oduct In	form	ation							
Pro	duct Typ	<u>م</u>	PLASMA	DERIVATIVE		ltem Code	(Source)			44-3944
	, uuce i yp	•					(000100)			
Da	ckaaina									
га. #	thom C	ada	Packa	ao Doscrinti	on	Markoting	Start Date	n Mai	katin	a End Data
# 1 N	IDC:0944-3	944-02		TON		Marketing			KELIII	g Lild Date
<b>•••</b>	antity o	f Dard	te la							
Qu	ancicy 0 + #	r Fait	ls Dackado (	Nuantity			Total Pro	duct Ou	antit	<b>V</b>
Parl	• <del>#</del> • 1 BOT	TIF	ackage	uantity		10 ml	Total Flo	uuct qu		у
Parl	t 2 1 VIAL	., GLASS	5			10 mL				
Pa	rt 1 of	f 2								
HEMOFIL M										
					0.010					
D	oduct I.	for	otion							
Pr	bauct In	orma								
lte	m Code (	Source	)	NDC:0944-3945	5					
Rou	ute of Ad	minist	ration	INTRAVENOUS						

Active Ingredier	nt/Active	Moiety						
	Ingre	edient Name		Basis of Str	ength	Strength		
ANTIHEMOPHILIC FA FACTOR HUMAN - UNII:	CTOR HUM 839MOZ 74GI	<b>AN</b> (UNII: 839MOZ74GK) (ANTIHEMOP <)	PHILIC	ANTIHEMOPHILI FACTOR HUMAN	C N	1000 [iU] in 10 mL		
Inactive Ingredi	ents							
		Ingredient Name			S	trength		
ALBUMIN HUMAN (UN	NII: ZIF514RV	ZR)						
HISTIDINE (UNII: 40D								
	5575672,							
Packaging								
# Item Code		Package Description		Marketi Start Da	ng ate	Marketing End Date		
<b>1</b> NDC:0944- 3945-01 10 mL Produc	in 1 BOTTLE; t (e.g., Drug/	Type 9: Other Type of Part 3 Combi Device/Biological Product)	ination					
Marketing In	Marketing Information							
Marketing Application Number or Monograph Marketing Start					Mark	ceting End		
Category		Citation	Date		Date			
BLA	BLA101448		02/23/19	88				
Part 2 of 2								
STERILE WAT	TER							
water liquid								
Product Inform	ation							
Item Code (Source	e)	NDC:64764-516						
Route of Administ	ration	INTRAVENOUS						
Inactive Ingredi	ents							
Ingredient Name				Stre	ngth			
WATER (UNII: 059QF0	KO0R)		10 ml	_ in 10 mL				
Packaging								
" Item		Deckare Description		Market	ing	Marketing		

# Co	ode		Package Descr	iption		Start Date End Da		End Date
<b>1</b> NDC:	64764- 10 m	L in 1 VIAL, GI	ASS; Type 9: Other T	ype of Part 3 C	Combination			
510-		uct (e.g., Drug	Device/Biological Fro	uuct)				
Marketing Information								
Ма	rketing	Applica	tion Number or M	onograph	Marketin	g Start	Ма	rketing End
Ca	ategory		Citation	5.	Dat	e		Date
BLA		BLA101448			02/23/1988			
Markating Information								
Ма		Annlica	tion Number or M	onograph	Marketin	a Start	Ma	rketing End
Ca	ategory	Аррпса	Citation	onograph	Dat	e	Ma	Date
BLA		BLA101448			02/23/1988			
LEM								
antiher	OFIL IVI	tor human	kit					
ununci								
Product Information								
Produ	ct Type	PLASMA	DERIVATIVE	ltem Code	(Source)	N	DC:094	44-3946
liouu	et type				(000100)			
Packa	aging							
# I	tem Code	Packa	ge Description	Marketing	ting Start Date N		Marketing End Date	
1 NDC:	0944-3946-02	2 1 in 1 CAI	RTON					
Quan	tity of Pa	rts						
Part #	:	Package (	Quantity		Total Pro	duct Qua	antity	1
Part 1	1 BOTTLE			10 mL				
Part 2	1 VIAL, GLA	SS		10 mL				
Dest	1 - 4 - 2							
Part	1 07 2							
HEM	OFIL M							
antihe	mophilic fa	ctor human	powder, for soluti	on				
Produ	uct Inform	nation						
ltem C	ode (Sour	ce)	NDC:0944-3947					
Route	of Adminis	tration	INTRAVENOUS					

Active Ingredier	nt/Active	Moiety						
	Ingre	edient Name		Basis of St	rength	Strength		
ANTIHEMOPHILIC FA FACTOR HUMAN - UNII:	CTOR HUM 839MOZ 74GI	<b>AN</b> (UNII: 839MOZ74GK) (ANTIHEMOP <)	PHILIC /	ANTIHEMOPHIL FACTOR HUMAI	IC N	1700 [iU] in 10 mL		
Inactive Ingredi	ents							
		Ingredient Name			S	trength		
ALBUMIN HUMAN (UNII: ZIF514RVZR)								
	397907L)							
Packaging								
# Item Code		Package Description		Marketi Start Da	ng ate	Marketing End Date		
<b>1</b> NDC:0944- 3947-01 10 mL Product	in 1 BOTTLE; t (e.g., Drug/	Type 9: Other Type of Part 3 Combi Device/Biological Product)	nation					
Marketing In	format	ion						
Marketing Category	Applicat	tion Number or Monograph Citation	Marketing Start Date		Marketing End Date			
BLA	BLA101448		02/23/198	8				
Part 2 of 2								
STERILE WA	TER							
Product Inform	ation							
Item Code (Source	e)	NDC:64764-516						
Route of Administ	ration	INTRAVENOUS						
Inactive Ingredi	ents							
Ingredient Name					ength			
WATER (UNII: 059QF0K00R) 10 mL in 10 mL								
Packaging								
# Item Code		Package Description		Market Start D	ing ate	Marketing End Date		
<b>1</b> NDC:64764- 516-10 Produce	in 1 VIAL, GL ct (e.g., Drug,	ASS; Type 9: Other Type of Part 3 C /Device/Biological Product)	ombinatior	1				

Marketing Information								
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date					
BLA	BLA101448	02/23/1988						
Marketing Ir	oformation							
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date					
BLA	BLA101448	02/23/1988						

Labeler - Takeda Pharmaceuticals America, Inc. (039997266)

Establishment					
Name	Address	ID/FEI	Business Operations		
BAXALTA US Inc.		085206634	MANUFACTURE(0944-3940, 0944-3942, 0944-3944, 0944-3946)		

# Establishment

Name	Address	ID/FEI	Business Operations
BAXALTA US Inc.		009471603	MANUFACTURE(0944-3940, 0944-3942, 0944-3944, 0944-3946)

Revised: 3/2025

Takeda Pharmaceuticals America, Inc.