

fibryga[®]
Fibrinogen (Human)

octapharma

Formulary Integration

A guide for adding fibryga
to your hospital's formulary.

Contents

SECTION	PAGE
<u>A. Adding Fibryga to Your Hospital's Formulary</u>	03
<u>B. Why Fibryga</u>	05
<u>C. Securing a P&T Committee Review</u>	07
<u>D. Preparing for P&T Committee Review</u>	09
<u>E. Proposal Development</u>	11
<u>F. Resources</u>	13

Adding Fibryga to Your Hospital's Formulary

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Please see Indications and Select Important Safety Information on page 4 and Important Safety Information throughout.

Page 03

SECTION A

Adding Fibryga to Your Hospital's Formulary

The First Step Toward Changing Your Hospital's Major Bleeding Management Protocols

Thank you for recognizing the need and advocating to improve major bleeding protocols within your hospital system through the use of fibrinogen concentrates like fibryga.

This guide is designed to help you navigate the process of proposing the addition of fibryga to your institution's formulary.

By following these steps, you will be equipped to make a compelling case to the Pharmacy & Therapeutics (P&T) Committee about the benefits and risks of fibryga and how it can help evolve the standard of care at your institution.

Note that all institutions have their own timelines and processes for adding products to formulary. Make sure to modify your approach to the specific needs within your organization.

INDICATIONS AND SELECT IMPORTANT SAFETY INFORMATION

Indications and Usage

Fibryga is a human fibrinogen concentrate indicated for fibrinogen supplementation in bleeding patients with acquired fibrinogen deficiency and in treatment of acute bleeding episodes in patients with congenital fibrinogen deficiency, including afibrinogenemia and hypofibrinogenemia.

Fibryga is not indicated for dysfibrinogenemia.

Contraindications

Fibryga is contraindicated in individuals who have manifested severe immediate hypersensitivity reactions, including anaphylaxis, to fibryga or its components (Sodium Citrate Dihydrate; Glycine; L-Arginine Hydrochloride).

Please see Important Safety Information throughout. Please see fibryga [full Prescribing Information](#).

Why Fibryga



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SECTION B

Why Fibryga

Indication¹:

Fibryga is a human fibrinogen concentrate (FC) indicated for:

- fibrinogen supplementation in bleeding patients with acquired fibrinogen deficiency
- treatment of acute bleeding episodes in patients with congenital fibrinogen deficiency, including afibrinogenemia and hypofibrinogenemia

Limitation of Use¹:

- Fibryga is not indicated for dysfibrinogenemia.

Details About Fibryga:



Precise amount of fibrinogen per vial (1 g/vial)



Virally inactivated and nanofiltered



Shelf stable at room temperature for up to 48 months



Supplied with a convenient transfer device featuring a protective vacuum seal (Octajet)



Rapidly reconstituted within 5 to 10 minutes

SELECT IMPORTANT SAFETY INFORMATION

Warnings and Precautions

Monitor patients for early signs of hypersensitivity or allergic reactions. If necessary, discontinue administration and institute appropriate treatment.

Thrombotic events have been reported in patients receiving fibryga. Treatment with human fibrinogen concentrate has been associated with thrombosis at target plasma fibrinogen levels that were below 150 mg/dL. The thrombotic risks may be greater when the target fibrinogen plasma level is 150 mg/dL. Weigh the benefits of administration versus the risks of thrombosis.

Fibryga is made from pooled human plasma. Products made from human plasma may contain infectious agents, e.g., viruses and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent.

Please see Important Safety Information throughout. Please see fibryga [full Prescribing Information](#).

Securing a P&T Committee Review



SECTION C

Securing a P&T Committee Review

The P&T Committee is a multidisciplinary team responsible for managing the formulary system in a hospital setting. This committee evaluates new drugs, reviews therapeutic guidelines, and develops policies related to drug use.

The following process has been implemented successfully in many top-tier institutions across the United States.²



Proposal Team

Recruit members for your team who will help you craft your narrative and help provide support during the committee review.

Key proposal team members include the following specialties:
anesthesiology, surgery, and pharmacy.



P&T Agenda

Identify your P&T Committee members, and follow the process required to secure a spot on an upcoming meeting agenda.



P&T Presentation

Engage your proposal team to help build your presentation, and to possibly attend and participate in your P&T meeting.

Preparing for P&T Committee Review

SECTION D

Preparing for P&T Committee Review

Fibryga Essentials

These resources will help you gain an understanding of fibryga, including the prescribing information, clinical profile, and clinical data.

Please click the below buttons to find the relevant information you may need.

[Product Website](#)

[AFD Clinical Study Publication 1](#)

[AFD Clinical Study Publication 2](#)

[Additional Resources](#)

[Prescribing Information](#)

[CFD Clinical Study Publication 1](#)

[CFD Clinical Study Publication 2](#)

[CFD Clinical Study Publication 3](#)

Please click [this link](#) to reach out to US Medical Affairs if you would like additional information about fibryga, including clinical studies.

A man with short brown hair and glasses, wearing a light blue button-down shirt, is shown in profile from the chest up. He is gesturing with his right hand as if speaking to a group of people. In the background, several other people are seated, some wearing white lab coats, suggesting a professional or medical setting. The background is blurred.

Proposal Development

SECTION E

Proposal Development

Ensure you are covering the following key topics when developing your proposal:

Highlight the unmet needs around fibrinogen replacement within your hospital setting for managing major bleed scenarios, especially where the current standard of care falls short.

Clearly identify the specific patient populations who are underserved by current protocols that utilize cryoprecipitate (cryo) to replace fibrinogen in major bleeding scenarios.

Demonstrate how fibrinogen concentrates (FC) like fibryga may improve patient outcomes when utilized in the management of major bleeding.²

Showcase the rapid administration and clinical benefits and risks of fibryga and how it may improve patient outcomes.²

Share case studies from your experience or from other institutions that are similar in location, size, level and experience that have successfully adopted fibryga.

SELECT IMPORTANT SAFETY INFORMATION

Adverse Reactions

The most serious adverse reactions observed with fibryga are thromboembolic episodes and anaphylactic-type reactions.

The most common adverse reactions observed in clinical studies with fibryga in acquired fibrinogen deficiency (>5% of patients) were abnormal hepatic function, acute kidney injury, anemia, atrial fibrillation, delirium and renal failure.

The most common adverse reactions observed in clinical studies with fibryga in congenital fibrinogen deficiency (>5% of patients) were nausea, vomiting, pyrexia (fever) and thrombocytosis.

Please see Important Safety Information throughout. Please see fibryga [full Prescribing Information](#).

Resources

Below are evidence-based content examples, based on specialty, to consider for your P&T Committee presentation.

They may help build your case around fibrinogen replacement in the context of major bleeding management and how fibrinogen concentrate (FC) can help address those needs by providing a different treatment approach.

To view specific content, simply click the appropriate link.

[Cardiac](#)

[Obstetrics](#)

[Trauma](#)

SECTION F

Cardiac Resources



Burden of major bleeding and low fibrinogen levels:

- Low plasma fibrinogen is an independent risk factor in perioperative bleeding.³
- Hypofibrinogenemia is observed in ~1/3 of cardiac surgery patients.⁴
- Up to 10% of cardiac surgeries are associated with severe or massive perioperative bleeding.⁵



Limitations of cryoprecipitate, the current standard of care:

- Cryoprecipitate (cryo) was initially developed to treat hemophilia A and von Willebrand disease. Its application has since expanded to fibrinogen replenishment in major bleeding scenarios.^{6,7}
- Despite rapid diagnosis via point-of-care testing, cryo availability is typically delayed by 25 to 45 minutes due to the need for ordering, thawing, and pooling from the blood bank.⁸⁻¹⁰
- Because cryo must be used within 4 to 6 hours of thawing, it leads to wastage rates of 7% to 33% and delays in treatment with thaw times of 15 to 30 minutes.^{8,11,12}



Clinical profile of fibrinogen concentrate:

- Recent updates to EU guidelines favor fibrinogen concentrate (FC) over cryo due to its standardization, ease of use, safety, and clinical efficacy.^{10,13-15}
- In cardiac procedures, FC use has been shown to decrease perioperative blood loss and transfusion requirements.^{16,17}
- The use of FC allows precise, consistent dosing, storage at room temperature, quick preparation without blood-type matching, and lower risk of infectious transmission through viral inactivation.^{3,18,19}
- The American Society of Anesthesiologists Guidelines suggest FC as an option in patients with excessive bleeding.²⁰

SELECT IMPORTANT SAFETY INFORMATION

Contraindications

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SECTION F

Obstetrics Resources



Burden of major bleeding and low fibrinogen levels:

- PPH accounts for about 14% of maternal deaths in the US.²¹
- Low fibrinogen levels are directly correlated with bleed severity during PPH.^{22,23}
- In pregnancy, fibrinogen levels below 200 mg/dL have a 100% positive predictive value for progressing to severe PPH.^{22,24}
- With each decrease of 100 mg/dL below 400 mg/dL the likelihood of severe bleeding is more than doubled.²²



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Clinical profile of fibrinogen concentrate:

- Use of fibrinogen concentrate (FC) swiftly restores fibrinogen levels and enhances maternal outcomes with minimal safety concerns.^{25,26}
- Availability of FC utilization for PPH resulted in faster administration of fibrinogen, and decreased utilization of packed red blood cell (pRBC) and fresh frozen plasma (FFP) transfusions compared to cryo.²⁷

SELECT IMPORTANT SAFETY INFORMATION

Warnings and Precautions

Monitor patients for early signs of hypersensitivity or allergic reactions. If necessary, discontinue administration and institute appropriate treatment.

Thrombotic events have been reported in patients receiving fibryga. Treatment with human fibrinogen concentrate has been associated with thrombosis at target plasma fibrinogen levels that were below 150 mg/dL. The thrombotic risks may be greater when the target fibrinogen plasma level is 150 mg/dL. Weigh the benefits of administration versus the risks of thrombosis.

Please see Important Safety Information throughout. Please see fibryga [full Prescribing Information](#).

SECTION F

Trauma Resources



Burden of major bleeding and low fibrinogen levels:

- Fibrinogen is the first coagulation factor to drop to critically low levels during bleeding, with levels of 150 to 200 mg/dL significantly increasing the risk of severe bleeding.^{3,7,28,29}
- Following acute trauma, fibrinogen levels below 150 mg/dL are linked to a significantly higher 28-day mortality rate (odds ratio [OR] 4.9).³⁰
- There is an ~4 times higher mortality rate in trauma patients with coagulopathy versus those without.³¹



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- Because cryo must be used within 4 to 6 hours of thawing, it leads to wastage rates of 7% to 33% and delays in treatment with thaw times of 15 to 30 minutes.^{8,11,12}



Clinical profile of fibrinogen concentrate:

- Fibrinogen concentrate (FC) use during trauma has demonstrated a reduction in blood loss and transfusion vs both cryo and control (did not receive FC within one hour after emergency admission).³²⁻³⁴

SELECT IMPORTANT SAFETY INFORMATION

Warnings and Precautions (cont.)

Fibryga is made from pooled human plasma. Products made from human plasma may contain infectious agents, e.g., viruses and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent.

Adverse Reactions

The most serious adverse reactions observed with fibryga are thromboembolic episodes and anaphylactic-type reactions.

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SECTION F

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