

**FUL-GLO- fluorescein sodium strip**  
**Akorn, Inc.**

*Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click [here](#).*

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**Fluorescein Sodium Ophthalmic Strips, USP**

Diagnostic Agent For Professional Use Only

Each sterile strip is impregnated with 0.6 mg of fluorescein sodium USP.

**INDICATIONS:**

For staining the anterior segment of the eye when fitting contact lenses, in disclosing corneal injury and in applanation tonometry.

**DIRECTIONS FOR USE:**

To ensure full fluorescence and patient comfort, the Ful-Glo impregnated tip should be moistened before application. One or two drops of sterile irrigation solution should be used for this purpose. Touch conjunctiva or fornix as required with moistened tip. It is recommended that the patient blink several times after application.

NOTE: Contents may not be sterile if individual strip package has been damaged or previously opened. Store below 30°C.

**INSTRUCTIONS FOR OPENING STERILE STRIPS:**

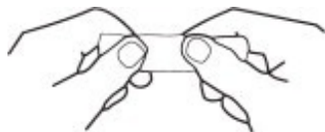
1. Separate tab ends. Pull apart slowly until white handle visible. Remove from envelope.



**Fig. 1**

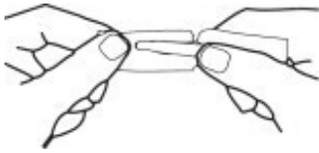
Or, for a convenient handle protector-

2. Grasp envelope firmly with two hands as shown in Fig. 2. Tear down the middle of envelope to strip.



**Fig. 2**

Hold the handle end of the strip firmly in the right hand as shown in Fig 3. Hold left edge of paper envelope so that tip is not held between the fingers.



**Fig. 3**

Snap the left portion off quickly. Strip is ready to use with a convenient handle protector.



**Fig. 4**

Principal Display Panel Text for Container Label:

NDC 17478-403-03 ONE STERILE STRIP

FUL-GLO® Fluorescein Sodium

Sterile Ophthalmic Strips 0.6 mg

Contents may not be sterile if this package has been damaged or previously opened. See box for complete information.

**NDC 17478-403-03** **ONE STERILE STRIP**  
**FUL-GLO®** **Fluorescein Sodium**  
**Sterile Ophthalmic Strips 0.6 mg**

**Contents may not be sterile if this package has been damaged or previously opened. See box for complete information.**

**Manufactured for Akorn, Inc., Lake Forest, IL 60045**

FGALP Rev. 09/09  
MSN 015-019

To  
Open,  
Pull Tab  
Apart  
Slowly



Principal Display Panel Text for Carton Label:

NDC 17478-403-03

FUL-GLO®

Fluorescein Sodium

Ophthalmic Strips, USP

0.6 mg

Diagnostic Agent for Professional Use Only

300 Sterile Strips

Akorn Logo



NDC 17478-403-03

**FUL-GLO<sup>®</sup>**  
**Fluorescein Sodium**  
**Ophthalmic Strips, USP**

0.6 mg

NDC 17478-403-03

**FUL-GLO<sup>®</sup>**  
**Fluorescein Sodium**  
**Ophthalmic Strips, USP**  
 0.6 mg



Lot No./Exp.

**FUL-GLO<sup>®</sup>**

**Fluorescein Sodium**  
**Ophthalmic Strips, USP**  
 Fluorescein Sodium Ophthalmic Strips, USP  
 Diagnostic Agent For Professional Use Only

Each sterile strip is impregnated with 0.6 mg of fluorescein sodium USP.

**INDICATIONS:** For staining the anterior segment of the eye when fitting contact lenses, in disclosing corneal injury and in appplanation tonometry.

**DIRECTIONS FOR USE:** To ensure full fluorescence and patient comfort, the Ful-Glo impregnated tip should be moistened before application. One or two drops of sterile irrigation solution should be used for this purpose. Touch conjunctiva or fornix as required with moistened tip. It is recommended that the patient blink several times after application.

**NOTE:** Contents may not be sterile if individual strip package has been damaged or previously opened. Store below 30°C.

**HOW SUPPLIED:** Dispenser carton containing 300 sterile strips.

**300 Sterile Strips**

MSN 015-021 FGALC Rev. 11/19  
 Manufactured for: **Akorn, Inc.**, Lake Forest, IL 60045

NDC 17478-403-03

**FUL-GLO<sup>®</sup>**  
**Fluorescein Sodium**  
**Ophthalmic Strips, USP**  
 0.6 mg

**INSTRUCTIONS FOR OPENING STERILE STRIPS:**

1. Separate tab ends. Pull apart slowly until white handle visible. Remove from envelope.



Fig. 1

**Or, for a conventional handle protector-**

2. Grasp envelope firmly with two hands as shown in Fig. 2. Tear down the middle of envelope to strip.

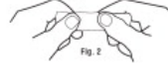


Fig. 2

Hold the handle end of the strip firmly in the right hand as shown in Fig. 3. Hold left edge of paper envelope so that tip is not held between the fingers.

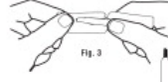


Fig. 3

Snap the left portion off quickly. Strip is ready to use with a conventional handle protector.



Fig. 4

**AKORN**

Diagnostic Agent For Professional Use Only  
 300 Sterile Strips

**FUL-GLO**

fluorescein sodium strip

**Product Information**

|  |  |  |                             |                           |
|--|--|--|-----------------------------|---------------------------|
| <b>Product Type</b>                    | HUMAN PRESCRIPTION DRUG  | <b>Item Code (Source)</b>                            | NDC:17478-403               |                           |
| <b>Route of Administration</b>         | OPHTHALMIC   |  |                             |                           |
| <b>Active Ingredient/Active Moiety</b> |  |  |                             |                           |
|  | <b>Ingredient Name</b>   | <b>Basis of Strength</b>                             | <b>Strength</b>             |                           |
|  | Fluorescein Sodium (UNI: 93X55PE38 X) (Fluorescein - UNI:TPY09G7XIR) | Fluorescein Sodium                                   | 0.6 mg                      |                           |
| <b>Packaging</b>                       |  |  |                             |                           |
| <b>#</b>                               | <b>Item Code</b>   | <b>Package Description</b>                           | <b>Marketing Start Date</b> | <b>Marketing End Date</b> |
| 1                                      | NDC:17478-403-03   | 300 in 1 CARTON                                      | 06/01/2004                  |                           |
| 1                                      |  | 1 in 1 APPLICATOR; Type 0: Not a Combination Product |                             |                           |
| <b>Marketing Information</b>           |  |  |                             |                           |
| <b>Marketing Category</b>              | <b>Application Number or Monograph Citation</b>                      | <b>Marketing Start Date</b>                          | <b>Marketing End Date</b>   |                           |
| unapproved drug other                  |  | 06/01/2004   |                             |                           |

**Labeler** - Akorn, Inc. (117696770)

**Registrant** - Akorn Operating Company LLC (117693100)

Revised: 10/2020

Akorn, Inc.