

# **CENTER FOR DRUG EVALUATION AND RESEARCH**

## **Approval Package for:**

***APPLICATION NUMBER:***

**NDA 20-287/S-034**

**Name:** Fragmin® (Dalteparin Sodium) Injection

**Sponsor:** Pharmacia & Upjohn

**Approval Date:** April 21, 2004

# CENTER FOR DRUG EVALUATION AND RESEARCH

***APPLICATION NUMBER:***

**NDA 20-287/S-034**

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# **CENTER FOR DRUG EVALUATION AND RESEARCH**

***APPLICATION NUMBER:***

**NDA 20-287/S-034**

**APPROVAL LETTER**



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration  
Rockville, MD 20857

NDA 20-287/S-034

Pharmacia & Upjohn Company  
Attention: Robert Clark  
Vice President, Regulatory Affairs  
235 E. 42<sup>nd</sup> Street  
New York, NY 10017

Dear Mr. Clark:

Please refer to your supplemental new drug application dated September 8, 2003, received September 9, 2003, submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Fragmin<sup>®</sup> (dalteparin sodium, injection).

We acknowledge receipt of your submission dated March 26, 2004.

Your submission of March 26, 2004, constituted a complete response to our March 9, 2004 action letter.

This "Changes Being Effected" supplemental new drug application provides for revisions to the DOSAGE AND ADMINISTRATION section of the package insert to add instructions to expel the air bubble prior to using the 10,000 IU single-dose graduated prefilled syringe.

We completed our review of this supplemental new drug application, as amended. It is approved, effective on the date of this letter, for use as recommended in the final printed labeling (FPL) submitted on March 26, 2004.

If you issue a letter communicating important information about this drug product (i.e., a "Dear Health Care Professional" letter), we request that you submit a copy of the letter to this NDA and a copy to the following address:

MEDWATCH, HFD-410  
FDA  
5600 Fishers Lane  
Rockville, MD 20857

We remind you that you must comply with the requirements for an approved NDA set forth under 21 CFR 314.80 and 314.81.

If you have any questions, call Diane Moore, Regulatory Project Manager, at (301) 827-7476.

Sincerely,

*{See appended electronic signature page}*

Robert L. Justice, M.D., M.S.

Director

Division of Gastrointestinal and Coagulation Drug

Products (HFD-180)

Office of Drug Evaluation III

Center for Drug Evaluation and Research

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/s/

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Joyce Korvick  
4/21/04 01:19:24 PM  
for Dr. Robert Justice

**CENTER FOR DRUG EVALUATION AND RESEARCH**

*APPLICATION NUMBER:*

**NDA 20-287/S-034**

**APPROVABLE LETTER**



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration  
Rockville, MD 20857

NDA 20-287/S-034

Pharmacia & Upjohn Company  
Attention: Gregory A. Brier,  
Senior Regulatory Manager  
Global Regulatory Affairs  
7000 Portage Road  
Kalamazoo, MI 49001

Dear Mr. Brier:

Please refer to your supplemental new drug application dated September 8, 2003, received September 10, 2003, submitted under section 505 of the Federal Food, Drug, and Cosmetic Act for Fragmin® (dalteparin sodium injection).

This "Changes Being Effected" supplemental new drug application provides for revising the instructions to expel the air bubble prior to using the graduated syringe.

We completed our review of this application, and it is approvable. Before this application may be approved, however, you must submit final printed labeling revised as follows:

1. In the **DOSAGE AND ADMINISTRATION** section of the package insert (PI), in the **Administration** subsection, *Instructions for using the prefilled single-dose syringes preassembled with passive needle guard devices* sub-subsection, in the fifth sentence that begins "Depress the plunger . . ." delete the phrase "To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection." so that the sentence reads "Depress the plunger of the syringe while holding the finger flange until the entire dose has been given."
2. In the **DOSAGE AND ADMINISTRATION** section, in the **Administration** subsection, *Graduated syringes* sub-sub-subsection, in the third sentence that reads "With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to depress the plunger down to the desired dose or volume, discarding the extra solution in an appropriate manner." delete the word "down" after the word "plunger" so that the sentence reads "With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to depress the plunger to the desired dose or volume, discarding the extra solution in an appropriate manner."
3. All previous revisions, as reflected in the most recently approved package insert, specifically S-032, must be included. To facilitate review of your submission, provide a highlighted or marked-up copy that shows the changes.

Please submit the final printed labeling (FPL) electronically according to the guidance for industry titled Providing Regulatory Submissions in Electronic Format – NDA. Alternatively, you may submit 20 paper copies of the FPL, as soon as it is available but no more than 30 days after it is printed. Please individually mount 15 of the copies on heavy-weight paper or similar material.

If additional information relating to the safety or effectiveness of this drug becomes available, revision of the labeling may be required.

Within 10 days after the date of this letter, you are required to amend this application, notify us of your intent to file an amendment, or follow one of your other options under 21 CFR 314.110. If you do not follow one of these options, we will consider your lack of response a request to withdraw the application under 21 CFR 314.65. Any amendment should respond to all the deficiencies listed. We will not process a partial reply as a major amendment nor will the review clock be reactivated until all deficiencies have been addressed.

If you have any questions, call Diane Moore, Regulatory Project Manager, at (301) 827-7476.

Sincerely,

*{See appended electronic signature page}*

Robert L. Justice, M.D., M.S.  
Director  
Division of Gastrointestinal & Coagulation Drug  
Products  
Office of Drug Evaluation III  
Center for Drug Evaluation and Research

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/s/

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Joyce Korvick  
3/9/04 09:52:35 AM  
for Dr. Robert Justice

# **CENTER FOR DRUG EVALUATION AND RESEARCH**

***APPLICATION NUMBER:***

**NDA 20-287/S-034**

**LABELING**





**CENTER FOR DRUG EVALUATION AND RESEARCH**

***APPLICATION NUMBER:***  
**NDA 20-287/S-034**

**LABELING REVIEWS**

**REGULATORY PROJECT MANAGEMENT LABELING**  
**Division of Gastrointestinal and Coagulation Drug Products**  
**(DGICDP)**

**Application Number:** NDA 20-287/SLR-034

**Name of Drug:** Fragmin® (dalteparin sodium, injection)

**Sponsor:** Pharmacia & Upjohn Company (a subsidiary of Pfizer)

**Materials Reviewed:** Package Insert (PI)

**Submission Date:** September 8, 2003

**Receipt Date:** September 9, 2003

**Background and Summary**

Fragmin is a low molecular weight heparin (LMWH) approved December 22, 1994, for use in the prophylaxis of deep venous thrombosis (DVT) which may lead to pulmonary embolism (PE) in patients undergoing hip replacement surgery and in patients undergoing abdominal surgery who are at risk for thromboembolic complications and for treatment of unstable angina and non-Q-wave myocardial infarction.

Labeling Supplement-031 (S-031) was submitted on January 14, 2003 (received January 15, 2003; approved on draft June 30, 2003) as a "Changes Being Effected" (CBE-O) supplement for the use of UltraSafe™ Passive needle safety guards in conjunction with the approved FRAGMIN® (dalteparin sodium injection) 10,000 IU/1.0 mL graduated pre-filled syringes. Among other sections revised in that labeling, the sub-subsection entitled "*Graduated syringes*" was added under the **Administration** subsection of the **DOSAGE AND ADMINISTRATION** section of the PI.

The most recently approved package insert (PI) for Fragmin is Efficacy Supplement-032 (S-032) (submitted February 7, 2003; received February 10, 2003; approved on draft December 10, 2003). S-032 is a prior approval efficacy supplement that added a new indication for the prophylaxis of deep vein thrombosis (DVT), which may lead to pulmonary embolism (PE) in medical patients who are at risk for thromboembolic complications due to restricted mobility during acute illness.

Supplement-034 is a Changes Being Effected (CBE) labeling supplement (submitted September 8, 2003; received September 9, 2003) that proposes to revise the instructions in the package insert (PI) that instruct the user to expel the air bubble prior to using the FRAGMIN graduated syringe.

Note: The revisions approved in S-032 (submitted February 7, 2003; received February 10, 2003; approved on draft December 10, 2003) were not included in the proposed labeling for S-034 (submitted September 8, 2003; received September 9, 2003) because S-034 was submitted three days before S-032 was approved.

### Review

The PI proposed for S-034 submitted September 8, 2003, received September 9, 2003, (identifying number 818 312-109) was compared to the FPL for S-032 (no identifying number) (submitted February 7, 2003; received February 10, 2003; approved on draft December 10, 2003). The proposed labeling for S-034 is identical to the approved labeling except for the following:

- I. The revisions made in the following sections in S-032 were not incorporated in the proposed text in the PI for S-034:

**CLINICAL TRIALS, INDICATIONS AND USAGE, WARNINGS, PRECAUTIONS, ADVERSE REACTIONS and DOSAGE AND ADMINISTRATION** sections.

The revisions should be included in the labeling to S-034. (See approval letter to S-032 dated December 10, 2003, and RPM Labeling review to S-032 dated November 6, 2003).

## II. **DOSAGE and ADMINISTRATION** section

### A. **Administration** subsection

1. In the *Subcutaneous injection technique* sub-subsection, the first paragraph that begins "Patients should be sitting . . ." the sponsor proposes to delete the second sentence that reads "to ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection."

This section pertains to the 10,000 IU and 25,000 IU graduated syringes. The sponsor explains in the cover letter that the air bubble should be expelled prior to discarding the extra solution in the 10,000 IU single-dose graduated syringe. Expelling the air bubble makes it easier to accurately determine the amount of solution that should be left in the syringe to obtain the desired dose. The sentence that had been added to this section in S-031 (submitted on January 14, 2003; received January 15, 2003; approved on draft June 30, 2003) to retain the air bubble applies only to the fixed-dose syringes (2500, 5000, 7500 IU syringes). The details for administering the fixed dose syringes and the graduated syringes are given in separate sections below the *Subcutaneous injection technique* section. The deletion of the sentence in this section avoids drawing the conclusion that the air bubble should not be expelled for

**subcutaneous injection. The deletion is acceptable per the Medical Officer, Dr. Ruyi He, in verbal communication to Diane Moore, RPM on February 3, 2004.**

2. *Instructions for using the prefilled single-dose syringes preassembled with passive needle guard devices* sub-subsection

- a. In the *Instructions for using the prefilled single-dose syringes preassembled with passive needle guard devices* sub-sub-subsection the sponsor changed the font from bold underlined letters to bold italicized letters.

**The revision is editorial and acceptable.**

b. *Fixed dose syringes* sub-sub-subsection

- 1) Before the first sentence, the sponsor inserted the sentence that reads "To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection."

**This is the same sentence that was added in S-031 (submitted January 14, 2003; received January 15, 2003; approved on draft June 30, 2003) in the *Subcutaneous injection technique* section. It fits more appropriately here to instruct the user to not expel the air bubble from the fixed-dose syringe (as opposed to the graduated syringes). The addition of the sentence to this section is acceptable per the Medical Officer, Dr. Ruyi He, in a verbal communication to Diane Moore, RPM on February 3, 2004.**

- 2) In the fifth sentence that begins, "Depress the plunger . . ." the sponsor has inserted the same sentence as above ("To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection.") in the middle of the phrase "until the entire dose has been given" so that the sentence reads as follows:

**"Depress the plunger of the syringe while holding the finger flange until the entire dose has To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection. been given."**

**This appears to be a typographical error as the new sentence does not belong in the middle of the original sentence. The revision is not acceptable.**

**Note: The revision appeared in the WORD version submitted September 8, 2003 (received September 9, 2003) but not in the PDF version submitted September 8, 2003 (received September 9, 2003); both versions were submitted together as a package. Both are identified as**

**“818 312 109”, however, the PDF version has an additional identification number of “5R6842 236.”**

**c. *Graduated syringes* sub-sub-subsection**

In the third sentence that reads “With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to depress the plunger to the desired dose or volume, discarding the extra solution in an appropriate manner.” the sponsor has replaced the word “depress” with the word “push” and added the word “down” after the word “plunger” so that the sentence reads “With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to push the plunger down to the desired dose or volume, discarding the extra solution in an appropriate manner.”

- 1) The replacement of the word “depress” with the word “push” is acceptable per the Medical Officer, Dr. Ruyi He, in verbal communication to Diane Moore, RPM on February 3, 2004.**
- 2) The inclusion of the word “down” does not make sense since the instructions tell the reader to hold the syringe pointing up. The word “down” should be deleted from the sentence.**

**Conclusions**

- 1. The revisions made in S-032 in the following item should be incorporated in the text of S-034: I.**
- 2. The following items are acceptable: II.A.2.a. and II.A.2.c. 1).**
- 3. The following items are acceptable per Dr. Ruyi He, Medical Officer: II.A.1. and II.A.2.b.1).**
- 4. The following items are not acceptable: II.A.2.b.2). and II.A.2.c.2).**
- 5. Because this supplement is a CBE supplement, the supplement should not be approved until the apparent typographical error in the DOSAGE AND ADMINISTRATION section, *Fixed dose syringes* sub-sub-subsection is corrected, the wording in the DOSAGE AND ADMINISTRATION section, *Graduated syringes* sub-sub-subsection is corrected and the revisions made in S-032 are incorporated into the labeling for S-034.**

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Diane Moore, B.S.  
Regulatory Health Project Manager

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Ruyi He, M.D.  
Medical Officer

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Kathy Robie-Suh, M.D., Ph.D.  
Hematology Team Leader

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Julieann DuBeau, MSN, RN  
Chief, Project Management Staff

Drafted: dm/January 15, 2004  
Revised: J.DuBeau 2.5.04/K.Robie-Suh 2.6.04  
Initialed: J.DuBeau 2.5.04/R.He, K.Robie-Suh 2.6.04  
Finalized: February 9, 2004  
Filename: N20287S34LbrevSV.doc

**RPM LABELING REVIEW**

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/s/

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Diane V. Moore  
2/6/04 03:29:59 PM  
CSO

Ruyi He  
2/6/04 05:01:50 PM  
MEDICAL OFFICER

Kathy Robie-Suh  
2/9/04 09:24:10 AM  
MEDICAL OFFICER

Julieann DuBeau  
2/9/04 12:06:59 PM  
CSO

**REGULATORY PROJECT MANAGEMENT LABELING**  
**Division of Gastrointestinal and Coagulation Drug Products**  
**(DGICDP)**

**Application Number:** NDA 20-287/SLR-034

**Name of Drug:** Fragmin® (dalteparin sodium, injection)

**Sponsor:** Pharmacia & Upjohn Company (a subsidiary of Pfizer)

**Materials Reviewed:** Package Insert (PI)

**Submission Date:** March 26, 2004

**Receipt Date:** March 29, 2004

**Background and Summary**

Fragmin is a low molecular weight heparin (LMWH) approved December 22, 1994, for use in the prophylaxis of deep venous thrombosis (DVT) which may lead to pulmonary embolism (PE) in patients undergoing hip replacement surgery and in patients undergoing abdominal surgery who are at risk for thromboembolic complications and for treatment of unstable angina and non-Q-wave myocardial infarction.

The most recently approved package insert (PI) for Fragmin is Efficacy Supplement-032 (S-032) (submitted February 7, 2003; received February 10, 2003; approved on draft December 10, 2003, no identifier code). S-032 is a prior approval efficacy supplement that added a new indication for the prophylaxis of deep vein thrombosis (DVT), which may lead to pulmonary embolism (PE) in medical patients who are at risk for thromboembolic complications due to restricted mobility during acute illness.

The sponsor submitted final printed labeling (FPL) for S-032 on January 9, 2004, (received January 10, 2004, identifier code "5R7065 376 818 312 111"). That PI included revisions proposed in S-034 and was found to be unacceptable (see RPM review to S-032 FPL by Diane Moore dated March 19, 2004.)

Supplement-034 is a Changes Being Effected (CBE) labeling supplement (submitted September 8, 2003; received September 9, 2003, identifier number "818 312 109") that proposes to revise the instructions in the PI that instruct the user to expel the air bubble prior to using the FRAGMIN graduated syringe. The revisions approved in S-032 (submitted February 7, 2003; received February 10, 2003; amended December 10, 2003, approved on draft December 10, 2003) were not included in the proposed labeling for S-034 (submitted September 8, 2003; received September 9, 2003) because S-034 was submitted three days before S-032 was approved. On March 9, 2004, DGICDP sent Pharmacia & Upjohn an approvable letter requesting the sponsor to 1) Correct the apparent typographical error in the **DOSAGE AND**

**ADMINISTRATION** section, **Administration** subsection, regarding deleting the phrase "To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection" from the middle of the sentence "Depress the plunger of the syringe while holding the finger flange until the entire dose has been given." 2) delete the word "down" after the word "plunger" in the third sentence of the **DOSAGE AND ADMINISTRATION** section, **Administration** subsection, *Graduated syringes* sub-sub-subsection so that the sentence reads "With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to depress the plunger to the desired dose or volume, discarding the extra solution in an appropriate manner." and 3) include all previous revisions, as reflected in the most recently approved package insert, specifically S-032 (submitted February 7, 2003; received February 10, 2003; amended December 10, 2003, approved on draft December 10, 2003).

The sponsor submitted revised FPL for S-034 on March 26, 2004 (received March 29, 2004).

### Review

The PI proposed for S-034 submitted March 26, 2004, received March 29, 2004, (identifying number 818 312-112B) was compared to the approved labeling for S-032 (no identifying number) (submitted February 7, 2003; received February 10, 2003; amended December 10, 2003, approved on draft December 10, 2003) and the approvable letter to S-034 dated December 10, 2003, with the list of deficiencies to S-034. The sponsor incorporated the revisions made in S-032 into the proposed PI text for S-034 (see RPM Labeling Review to S-034 dated February 9, 2004, by Diane Moore). The proposed labeling for S-034 is identical to the approved labeling in S-032 except for the following:

#### **I. DOSAGE and ADMINISTRATION section**

##### **A. Administration subsection**

1. In the *Subcutaneous injection technique* sub-subsection, the first paragraph that begins "Patients should be sitting . . ." the sponsor deleted the second sentence that reads "to ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection."

**The deletion is acceptable (see RPM Labeling Review to S-034 dated February 9, 2004, by Diane Moore).**

2. *Instructions for using the prefilled single-dose syringes preassembled with passive needle guard devices* sub-subsection
  - a. In the *Instructions for using the prefilled single-dose syringes preassembled with passive needle guard devices* sub-sub-subsection the sponsor changed the font from bold underlined letters to bold italicized letters.

**The revision is editorial and acceptable.**

b. *Fixed dose syringes* sub-sub-subsection

- 1) Before the first sentence, the sponsor inserted the sentence that reads "To ensure delivery of the full dose, do not expel the air bubble from the prefilled syringe before injection."

**The deletion is acceptable (see RPM Labeling Review to S-034 dated February 9, 2004, by Diane Moore).**

Note: In the fifth sentence that begins, "Depress the plunger . . ." the sponsor has corrected the typographical error from the sentence. (See RPM Labeling Review to S-034 dated February 9, 2004, by Diane Moore).

c. *Graduated syringes* sub-sub-subsection

In the third sentence that reads "With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to depress the plunger to the desired dose or volume, discarding the extra solution in an appropriate manner." the sponsor has replaced the word "depress" with the word "push" and deleted the previously added the word "down" after the word "plunger" so that the sentence reads "With the needle pointing up, prepare the syringe by expelling the air bubble and then continuing to push the plunger to the desired dose or volume, discarding the extra solution in an appropriate manner."

**The revisions are acceptable (See RPM Labeling Review to S-034 dated February 9, 2004, by Diane Moore).**

### **Conclusions**

1. The proposed revisions made to the FPL submitted to S-034 on March 26, 2004, received March 29, 2004, are acceptable.
2. The FPL to S-034 submitted March 26, 2004 (received March 29, 2004) should be approved.
3. Because FPL was submitted to S-034, this labeling supercedes the FPL to S-032. The FPL labeling to S-032 dated February 7, 2003, (received February 10, 2003) should be retained in the files, but not be referenced as approved FPL. The labeling submitted to S-034 on March 26, 2004, received March 29, 2004 is now the currently approved labeling for Fragmin NDA 20-287.

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Diane Moore, B.S.  
Regulatory Health Project Manager

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Julieann DuBeau, MSN, RN  
Chief, Project Management Staff

Drafted: dm/April 7, 2004  
Revised: J.DuBeau 4.14.04  
Initialed: J.DuBeau 4.14.04  
Finalized: April 15, 2004  
Filename: N20287S34Lblrev32904.doc

**RPM LABELING REVIEW**

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/s/

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Diane V. Moore  
4/15/04 12:52:15 PM  
CSO

Julieann DuBeau  
4/19/04 03:09:31 PM  
CSO

# **CENTER FOR DRUG EVALUATION AND RESEARCH**

*APPLICATION NUMBER:*

**NDA 20-287/S-034**

**ADMINISTRATIVE and CORRESPONDENCE**  
**DOCUMENTS**

**Pfizer Inc**  
7000 Portage Road  
Kalamazoo, MI 49001  
Tel 269.833.4000



September 8, 2003

Central Document Room  
Center for Drug Evaluation and Research  
Food and Drug Administration  
12229 Wilkins Avenue  
Rockville, MD 20852

**Re: NDA 20-287**  
**FRAGMIN®**  
**dalteparin sodium injection**

**ELECTRONIC SUBMISSION**

**Special Labeling Supplement**  
**Changes Being Effected (CBE)**  
**Graduated Syringe - Expel Air Bubble**

Dear Sir/Madam:

Pursuant to CFR 314.70(c)(2)(iii), Pfizer is submitting a "Special Labeling Supplement – Changes Being Effected," to NDA-20-287, FRAGMIN® (dalteparin sodium injection) revising instructions to expel the air bubble prior to using the graduated syringe.

Syringes pre-filled with FRAGMIN contain an air bubble as a result of the assembly process. In Supplement 20-287/S-031, we revised the DOSAGE AND ADMINISTRATION section of the package insert for FRAGMIN to instruct health care professionals that, to ensure that the full dose of FRAGMIN is delivered, they should not expel the air bubble from the pre-filled syringes. Supplement 031 was approved by the Agency on June 30, 2003. We have become aware that the instruction to retain the air bubble applies only to the fixed-dose syringes (2500, 5000, 7500 IU syringes). For the 10,000 IU single-dose graduated syringe, the air bubble should be expelled prior to discarding the extra solution in the syringe. Expelling the air bubble makes it easier to accurately determine the amount of solution that should be left in the syringe to obtain the desired dose. Therefore, this special labeling supplement provides for the correction of the air bubble instruction in the DOSAGE AND ADMINISTRATION section of the package insert.

Final printed labeling is being submitted electronically as follows:

Pharmacia & Upjohn Company is the Sponsor of NDA 20-287 FRAGMIN® (dalteparin sodium injection) and a wholly owned subsidiary of Pharmacia Corporation. Pharmacia Corporation is a wholly owned subsidiary of Pfizer. Pfizer is the authorized agent for this NDA.

- Package insert, identified by copy code 818 312 109 (electronic file named *pi.pdf*)
- Marked up changes are provided in a MS WORD version of the package insert (electronic file named *mockup.doc*)

Pfizer plans to implement the use of the revised package insert for FRAGMIN® no later than November 30, 2003.

The CD-ROM contains the following files and directory structure:

**Main Directory – N20287**

- Cover Letter (cover.pdf)
- 356h Form (356h.pdf)
- Table of Contents (ndatoc.pdf)

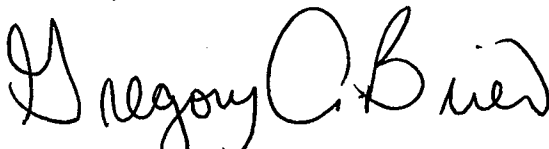
**Subdirectory – Labeling:**

- Final Printed Labeling, Package Insert copy code 818 312 109 (pi.pdf)
- Package Insert Marked Up in MS Word (mockup.doc)
- Labeling Table of Contents (labeltoc.pdf)

Each CD-ROM has been scanned with Trend Micro OfficeScan Corporate Edition for Windows NT version 5.02 and found to be virus free.

If you have questions regarding this correspondence, please contact either Ms. Alexandra Pearce, Regulatory Liaison Director, World Wide Strategy by telephone at 212-733-6079 or by fax at 212-857-3558 or myself by telephone at (269) 833-3670 or by fax at (269) 833-8237.

Sincerely,



Gregory A. Butler, BScHE, MBA  
Senior Regulatory Manager  
Global Regulatory Affairs

GAB:mlw



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration  
Rockville, MD 20857

NDA 20-287/S-034

**CBE-0 SUPPLEMENT**

Pharmacia & Upjohn Company  
Attention: Gregory A. Brier  
Senior Regulatory Manager  
7000 Portage Road, Kalamazoo, MI 49001

Dear Mr. Brier:

We have received your supplemental drug application submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for the following:

Name of Drug Product:      Fragmin (dalteparin sodium, injection)

NDA Number:                20-287

Supplement number:        S-034

Date of supplement:        September 8, 2003

Date of receipt:             September 9, 2003

This supplemental application, submitted as "Supplement - Changes Being Effected" proposes the following change: Revision of the **DOSAGE AND ADMINISTRATION** section in the package insert to add instructions to expel the air bubble prior to using the 10,000 IU single-dose graduated prefilled syringe.

We filed the application on November 8, 2003, in accordance with 21 CFR 314.101(a). The user fee goal date is March 9, 2004.

All communications concerning this supplement should be addressed as follows:

U.S. Postal Service:

Center for Drug Evaluation and Research  
Division of Gastrointestinal and Coagulation Drug Products HFD-180  
Attention: Division Document Room, 8B-45  
5600 Fishers Lane  
Rockville, Maryland 20857

Courier/Overnight Mail:

Food and Drug Administration

Center for Drug Evaluation and Research

Division of Gastrointestinal and Coagulation Drug Products HFD-180

Attention: Division Document Room, 8B-45

5600 Fishers Lane

Rockville, Maryland 20857

If you have any questions, call me at (301) 827-7476.

Sincerely,

*{See appended electronic signature page}*

Diane Moore

Regulatory Project Manager

Division of Gastrointestinal and Coagulation

Drug Products HFD-180

Office of Drug Evaluation III

Center for Drug Evaluation and Research

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/s/

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Diane V. Moore  
11/20/03 05:41:18 PM