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New Drug Application (NDA): 020287  
Company: PFIZER INC

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| Products on NDA 020287 |                    |                              |                         |
|------------------------|--------------------|------------------------------|-------------------------|
| CSV                    | Excel              | Print                        |                         |
| Drug Name              | Active Ingredients | Strength                     | Dosage Form/Route       |
| + FRAGMIN              | DALTEPARIN SODIUM  | 2,500IU/0.2ML (12,500IU/ML)  | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 10,000IU/0.4ML (25,000IU/ML) | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 5,000IU/0.2ML (25,000IU/ML)  | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 10,000IU/ML (10,000IU/ML)    | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 7,500IU/0.3ML (25,000IU/ML)  | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 95,000IU/3.8ML (25,000IU/ML) | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 95,000IU/9.5ML (10,000IU/ML) | INJECTABLE;SUBCUTANEOUS |
| + FRAGMIN              | DALTEPARIN SODIUM  | 7,500 IU/0.75ML              | INJECTABLE;INJECTION    |

|                                                                                           |                   |                               |                         |
|-------------------------------------------------------------------------------------------|-------------------|-------------------------------|-------------------------|
|  FRAGMIN  | DALTEPARIN SODIUM | 12,500IU/0.5ML (25,000IU/ML)  | INJECTABLE;SUBCUTANEOUS |
|  FRAGMIN | DALTEPARIN SODIUM | 15,000IU/0.6ML (25,000IU/ML)  | INJECTABLE;SUBCUTANEOUS |
|  FRAGMIN | DALTEPARIN SODIUM | 18,000IU/0.72ML (25,000IU/ML) | INJECTABLE;SUBCUTANEOUS |

Showing 1 to 11 of 11 entries

Approval Date(s) and History, Letters, Labels, Reviews for NDA 020287 

Labels for NDA 020287 

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