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New Drug Application (NDA): 007073
Company: PHARMACIA AND UPJOHN

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Products on NDA 007073

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Drug Name	Active Ingredients	Strength	Dosage Form/Route	Marketing Status
AZULFIDINE	SULFASALAZINE	500MG	TABLET;ORAL	Prescription
AZULFIDINE EN-TABS	SULFASALAZINE	500MG	TABLET, DELAYED RELEASE;ORAL	Prescription

Showing 1 to 2 of 2 entries

Approval Date(s) and History, Letters, Labels, Reviews for NDA 007073

Labels for NDA 007073

Therapeutic Equivalents for NDA 007073

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U.S. Food and Drug Administration

10903 New Hampshire Avenue
Silver Spring, MD 20993
1-888-INFO-FDA (1-888-463-6332)

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