



FDA Approved Drug Products

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Label and Approval History

Drug Name(s) DEPO-MEDROL (Brand Name Drug)
 FDA Application No. (NDA) 011757
 Active Ingredient(s) METHYLPREDNISOLONE ACETATE
 Company PHARMACIA AND UPJOHN

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Label Information

What information does a label include?

Note: Not all labels are available in electronic format from FDA.

Labels are not available on this site for DEPO-MEDROL, NDA no. 011757

Approval History NDA 011757

Note: Not all reviews are available in electronic format from FDA.

Older labels are for historical information only, and should not be used for clinical purposes.
Action dates can only be verified from 1984 to the present.

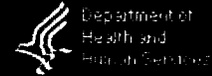
Click on a column header to re-sort the table:

Action Date	Supplement Number	Approval Type	Letters, Reviews, Labels, Patient Package Insert	Note
05/22/2002	079	Control Supplement		This supplement type does not usually require new labeling.
04/04/2000	078	Manufacturing Change or Addition		This supplement type does not usually require new labeling.
01/24/1995	075	Package Change		Label is not available on this site.
01/07/1994	072	Manufacturing Change or Addition		This supplement type does not usually require new labeling.
12/28/1993	070	Package Change		Label is not available on this site.
11/15/1990	064	Formulation Revision		Label is not available on this site.
12/14/1989	063	Labeling Revision		Label is not available on this site.
01/05/1989	059	Package Change		Label is not available on this site.
05/12/1986	053	Labeling Revision		Label is not available

"Adhesive Arachnoiditis"

Please read about this debilitating condition before having a spinal procedure done. The neurotoxin damage done to the central nervous system is permanent and the pain is continuous. There is no cure or dedicated research. Some invasive spinal procedures that can result in Arachnoiditis are: Epidurals for child birth, Spinal injections, Spinal diagnostics, and Surgery. Personal research (Internet search) can save a lifetime of suffering for yourself or someone you know. This risk is rarely disclosed.

Website: www.cofwa.org



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Therapeutic Equivalents

Drug Name(s) METHYLPREDNISOLONE ACETATE (Generic Drug)

FDA Application No. (ANDA) 040620

Active Ingredient(s) METHYLPREDNISOLONE ACETATE

Company SICOR PHARMS

2nd Print

INJECTABLE; INJECTION; 40MG/ML

TE Code = AB

Drug Name	Active Ingredients	Strength	Dosage Form/Route	Marketing Status	RLD	TE Code	Application Number	Company
DEPO-MEDROL	METHYLPREDNISOLONE ACETATE	40MG/ML	INJECTABLE; INJECTION	Prescription	Yes	AB	011757	PHARMACIA AND UPJOHN
METHYLPREDNISOLONE ACETATE	METHYLPREDNISOLONE ACETATE	40MG/ML	INJECTABLE; INJECTION	Prescription	No	AB	040557	SICOR PHARMS
METHYLPREDNISOLONE ACETATE	METHYLPREDNISOLONE ACETATE	40MG/ML	INJECTABLE; INJECTION	Prescription	No	AB	040620	SICOR PHARMS

INJECTABLE; INJECTION; 80MG/ML

TE Code = AB

Drug Name	Active Ingredients	Strength	Dosage Form/Route	Marketing Status	RLD	TE Code	Application Number	Company
DEPO-MEDROL	METHYLPREDNISOLONE ACETATE	80MG/ML	INJECTABLE; INJECTION	Prescription	Yes	AB	011757	PHARMACIA AND UPJOHN
METHYLPREDNISOLONE ACETATE	METHYLPREDNISOLONE ACETATE	80MG/ML	INJECTABLE; INJECTION	Prescription	No	AB	040557	SICOR PHARMS
METHYLPREDNISOLONE ACETATE	METHYLPREDNISOLONE ACETATE	80MG/ML	INJECTABLE; INJECTION	Prescription	No	AB	040620	SICOR PHARMS

• Labels are not available

• Approval History, Letters, Reviews, and Related Documents

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2004 - 2005, approved

"Adhesive Arachnoiditis"

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Drug Details

Drug Name(s) DEPO-MEDROL (Brand Name Drug)
FDA Application No. (NDA) 011757
Active Ingredient(s) METHYLPREDNISOLONE ACETATE
Company PHARMACIA AND UPJOHN
Original Approval or Tentative Approval Date May 27, 1959
Chemical Type 2 New ester, new salt, or other noncovalent derivative
Review Classification S Standard review drug

Most recent part of this

- Therapeutic Equivalents
- Approval History, Letters, Reviews, and Related Documents
- Labels are not available

this was just added to Drugs@FDA website

Products on Application (NDA) #011757

Click on a column header to re-sort the table:

Drug Name	Active Ingredients	Strength	Dosage Form/Route	Marketing Status	RLD	TE Code
DEPO-MEDROL	METHYLPREDNISOLONE ACETATE	40MG/ML	INJECTABLE; INJECTION	Prescription	Yes	AB
DEPO-MEDROL	METHYLPREDNISOLONE ACETATE	20MG/ML	INJECTABLE; INJECTION	Prescription	Yes	None
DEPO-MEDROL	METHYLPREDNISOLONE ACETATE	80MG/ML	INJECTABLE; INJECTION	Prescription	Yes	AB

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