



Center for Medicaid and State Operations

November 25, 2008

TO: Drug Rebate Technical Contacts

FROM: Medicaid Drug Rebate Program

SUBJECT: Deleted Product-Immediate Action Required

The FDA has determined that the following active NDC is an unapproved new drug within the meaning of section 201(p) of the Federal Food, Drug, and Cosmetic Act, subject to enforcement action, and cannot be marketed without appropriate FDA approval. February 8, 2008 (73 Fed. Reg. 7565). According to the FDA, this product does not have an approved New Drug Application; therefore, the NDC does not meet the definition of a covered outpatient drug as defined in Section 1927(k) of the Social Security Act and is subsequently no longer eligible for inclusion in the rebate program. **Consequently, this NDC should be deleted from your state Medicaid Drug Rebate system as of the date of this notice.** The labeler of this product is responsible for paying rebates on this NDC if it was dispensed prior to the date of this notice. In addition, states should be aware that the fourth quarter 2008 tape to states will be the last quarterly tape that will include this NDC in order to facilitate rebate billing for any utilization that occurred in good faith prior to the date of this notice. However, states are reminded that no Federal Financial Participation will be available for this drug after the date of this notice for purposes of the Medicaid Drug Rebate Program.

NDC	Product Name
55390-0605	COLCHICINE INJECTION USP