

**Request for Prior Authorization
SELECTED BRAND NAME DRUGS**

Iowa Medicaid MedWatch Form

Revised for submission of brand medically necessary requests for the Iowa Medicaid Pharmacy Program. Prescriber must have witnessed or has documentation that the manifestation of adverse event(s) is linked to generic drug. Completion of form does not automatically grant approval; incomplete forms will be returned with denial.***

A. PATIENT INFORMATION

Name: _____ Sex: F M
 Medicaid ID: _____ DOB: ____/____/____
 Weight: _____ lbs Phone: (____) _____
 Has a generic been tried before? Yes No

Give date: ____/____/____ Age at time of event: _____

1. Check all that apply Adverse
- Event Product
 - Use Error
 - Product Problem (e.g., defects/malfunctions)
 - Problem with Different Manufacturer of Same Medicine

2. Outcomes Attributed to Adverse Event: (Check all that apply.) Death:
- (month/day/year) Disability or Permanent Damage
 - Life-threatening
 - Congenital Anomaly/ Birth Defects
 - Required Intervention to Prevent Permanent Impairment/Damage
 - Hospitalization – Initial or Prolonged
 - Other Serious (Important Medical Events)

3. Date of Event (mo/day/yr) _____ 4. Date of This Report (mo/day/yr) _____

5. Describe Event, Problem, or Product Use Error; Relevant History & Tests

8. NDC # (specify generic manufacturer #1 _____ #2 _____)

Product names and therapy dates (exclude treatment of event)

Signature certifies that brand is medically necessary

Prescriber's Name _____
 Signature _____ NPI # _____
 Address: _____
 Phone #: (____) _____ - _____

Fax #: (____) _____ - _____

Did the prescriber witness the ADR? Yes No
 Has the ADR been previously reported to the FDA? Yes No

**Please FAX form to the Iowa Medicaid Pharmacy
 Programat
 1-877-733-3195
 DO NOT fax directly to the FDA**

1. Name (Give labeled strength & mfr/labeler, if known)

#1 _____
 #2 _____

2. Dose, Frequency & Route Used	3. Therapy Dates
#1 _____	#1 _____
#2 _____	#2 _____

4. Diagnosis for Use (Indication)	5. Event Abated After Use Stopped or Dose Reduced?
#1 _____	#1 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A
#2 _____	#2 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A

6. Lot # (if known)	7. Event Reappeared After Reintroduction
#1 _____	#1 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A
#2 _____	#2 <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A