

1998: Actiq Approved Indication

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3 **Actiq**[®]
4 (oral transmucosal fentanyl citrate)
5 CII
6
7 **PHYSICIANS AND OTHER HEALTHCARE PROVIDERS**
8 **MUST BECOME FAMILIAR WITH THE IMPORTANT**
9 **WARNINGS IN THIS LABEL.**
10
11 *Actiq* is indicated only for the management of breakthrough
12 cancer pain in patients with malignancies who are already
13 receiving and who are tolerant to opioid therapy for their
14 underlying persistent cancer pain. Patients considered opioid tolerant
15 are those who are taking at least 60 mg morphine/day, 50 µg transdermal
16 fentanyl/hour, or an equianalgesic dose of another opioid for a week or
17 longer.
18
19 Because life-threatening hypoventilation could occur at any dose in
20 patients not taking chronic opiates, *Actiq* is contraindicated in the
21 management of acute or postoperative pain. This product **must not** be
22 used in opioid non-tolerant patients.
23
24 *Actiq* is intended to be used only in the care of cancer patients and only
25 by oncologists and pain specialists who are knowledgeable and skilled in
26 the use of Schedule II opioids to treat cancer pain.
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28 Patients and their caregivers must be instructed that *Actiq* contains
29 a medicine in an amount which can be fatal to a child. Patients and
30 their caregivers must be instructed to keep all units out of the reach
31 of children and to discard opened units properly. (See information
32 for Patients and Their Caregivers for disposal instructions.)
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36 **WARNING:** May be habit forming
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40 **DESCRIPTION**
41 *Actiq* (oral transmucosal fentanyl citrate) is a solid formulation of
42 fentanyl citrate, a potent opioid analgesic, intended for oral transmucosal
43 administration. *Actiq* is formulated as a white to off-white solid drug
44 matrix on a handle that is radiopaque and is fracture resistant (ABS
plastic) under normal conditions when used as directed.

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PLAINTIFFS TRIAL
EXHIBIT
P-03606_00001