AN ACT
RELATING TO OPIOID OVERDOSE; REQUIRING HEALTH CARE PROVIDERS,
UNDER CERTAIN CIRCUMSTANCES, TO ADVISE PATIENTS ON THE RISKS
OF OVERDOSE AND OPIOID OVERDOSE REVERSAL MEDICATION AND TO
CO-PRESCRIBE AN OPIOID ANTAGONIST.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF NEW MEXICO:

SECTION 1. Section 24-2D-1 NMSA 1978 (being Laws 1999,
Chapter 126, Section 1) is amended to read:

"24-2D-1. SHORT TITLE.--Chapter 24, Article 2D
NMSA 1978 may be cited as the "Pain Relief Act"."

SECTION 2. Section 24-2D-2 NMSA 1978 (being Laws 1999,
Chapter 126, Section 2, as amended) is amended to read:

"24-2D-2. DEFINITIONS.--As used in the Pain Relief Act:

A. "accepted guideline" means the most current
clinical pain management guideline developed by the American
geriatrics society or the American pain society or a clinical
pain management guideline based on evidence and expert
opinion that has been accepted by the New Mexico medical
board;

B. "acute pain" means the normal, predicted
physiological response to a noxious chemical or thermal or
mechanical stimulus, typically associated with invasive
procedures, trauma or disease and generally time-limited;

C. "board" means the licensing board of a health
care provider;

D. "chronic pain" means pain that persists after reasonable medical efforts have been made to relieve the pain or its cause and that continues, either continuously or episodically, for longer than three consecutive months. "Chronic pain" does not include pain associated with a terminal condition or with a progressive disease that, in the normal course of progression, may reasonably be expected to result in a terminal condition;

E. "clinical expert" means a person who by reason of specialized education or substantial relevant experience in pain management has knowledge regarding current standards, practices and guidelines;

F. "disciplinary action" means any formal action taken by a board against a health care provider, upon a finding of probable cause that the health care provider has engaged in conduct that violates the board's practice act;

G. "health care provider" means a person who is licensed or otherwise authorized by law to provide health care in the ordinary course of business or practice of the person's profession and who has prescriptive authority within the limits of the person's license;

H. "opioid analgesic" means buprenorphine, butorphanol, codeine, hydrocodone, hydromorphine, levorphanol, meperidine, methadone, morphine, nalbuphine,
oxycodone, oxymorphone, pentazocine and propoxyphene as well as their brand names, isomers and combinations;

I. "opioid antagonist" means a drug approved by the federal food and drug administration that when administered negates or neutralizes in whole or in part the pharmacological effects of an opioid analgesic in the body, including naloxone and such other medications approved by the board of pharmacy for the reversal of opioid analgesic overdoses;

J. "pain" means acute and chronic pain; and

K. "therapeutic purpose" means the use of pharmaceutical and non-pharmaceutical medical treatment that conforms substantially to accepted guidelines for pain management."

SECTION 3. A new section of the Pain Relief Act is enacted to read:

"REQUIREMENTS FOR HEALTH CARE PROVIDERS WHO PRESCRIBE, DISTRIBUTE OR DISPENSE OPIOID ANALGESICS.--

A. A health care provider who prescribes, distributes or dispenses an opioid analgesic for the first time to a patient shall advise the patient on the risks of overdose and inform the patient of the availability of an opioid antagonist. With respect to a patient to whom an opioid analgesic has previously been prescribed, distributed or dispensed by the health care provider, the health care provider shall advise the patient on the risks of overdose and the availability of an opioid antagonist."
provider shall advise the patient on the risks of overdose and inform the patient of the availability of an opioid antagonist on the first occasion that the health care provider prescribes, distributes or dispenses an opioid analgesic each calendar year.

B. A health care provider who prescribes an opioid analgesic for a patient shall co-prescribe an opioid antagonist if the amount of opioid analgesic being prescribed is at least a five-day supply. The prescription for the opioid antagonist shall be accompanied by written information regarding the temporary effects of the opioid antagonist and techniques for administering the opioid antagonist. That written information shall contain a warning that a person administering the opioid antagonist should call 911 immediately after administering the opioid antagonist."