August 10, 2001

BUREAU CIRCULAR
No. ___ s. 2001

To : ALL DRUG MANUFACTURERS
Subject : Submission of Site Information File and updating.

To facilitate the evaluation and inspection of drug manufacturer's compliance to current Good manufacturing Practice (cGMP) and other related regulations, you are hereby directed to immediately submit your current site information file on or before October 15, 2001.

Changes in the information previously provided must be updated and submitted to Regulation Division II on or before every 15th of December.

[Signature]
WILLIAM D. TORRES
Director

SMOKING IS DANGEROUS TO YOUR HEALTH