## Drug Status Report

**Drug:** Phenobarbital related substances

2,4-Dimino-6-one phenobarbital (I)

4-Imino-2,6-diketo phenobarbital (II)

Methyl ethyl phenyl cyanoacetate (III)

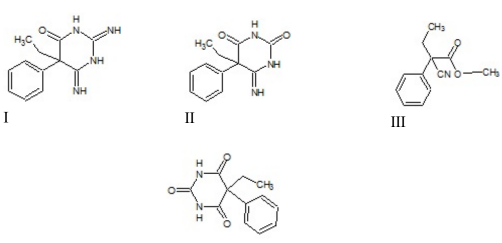
**Drug Name Status**: The above names are common names.

Chemical Name: (I) 5-Ethyl-5-phenyl-2,4-diamino-(5H)-pyrimidine-6-one

(II) 5-Ethyl-5-phenyl-4-diamino-(5H)-pyrimidine-2,6-dione

(III) Methyl 2-cyano-2-phenyl-butanoate

## **Chemical structure:**



Phenobarbital

Molecular Formula: (I) C<sub>12</sub>H<sub>14</sub>N<sub>4</sub>O

(II)  $C_{12}H_{13}N_3O_2$ (III)  $C_{12}H_{13}NO_2$ 

Pharmacological class / Application: pharmaceutical related substance

## **International status:**

US: The substances are not listed on the schedules to the CSA and are not mentioned on the DEA website.

United Nations: The substances are not listed on the Yellow List - List of Narcotic Drugs under International Control nor the Green List - List of Psychotropic Substances under International

## Control.

Canadian Status: None of substances I to III is not listed on the schedules to the CDSA. They are structurally similar to phenobarbital. Chemically, they are not barbiturates. Substances I to III are precursors in the synthesis of phenobarbital according to the scheme below.

Recommendation: 2,4-Dimino-6-one phenobarbital (I), 4-imino-2,6-diketo phenobarbital (II) and methyl ethyl phenyl cyanoacetate (III) are not included in the schedules to the CDSA and are not controlled substances.

February 5, 2009