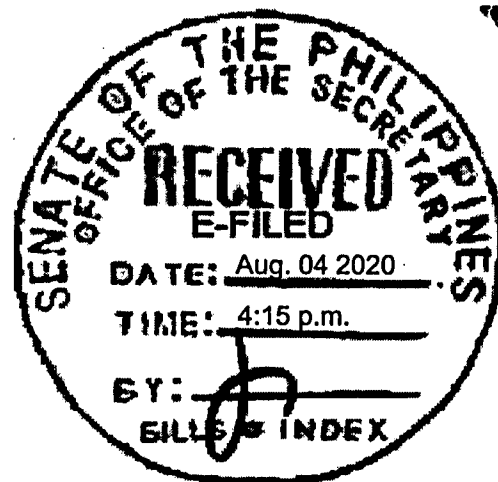


EIGHTEENTH CONGRESS OF THE )  
REPUBLIC OF THE PHILIPPINES )  
Second Regular Session )

SENATE

S.B. No. 1760



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Introduced by **SEN. IMEE R. MARCOS**

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**AN ACT ESTABLISHING THE DRUG PRICE REGULATORY BOARD TO  
REGULATE THE PRICES OF DRUGS AND MEDICINES IN THE PHILIPPINES  
AMENDING FOR THE PURPOSE REPUBLIC ACT NO. 9502, OTHERWISE  
KNOWN AS THE "UNIVERSALLY ACCESSIBLE CHEAPER AND QUALITY  
MEDICINES ACT OF 2008" AND FOR OTHER PURPOSES**

**EXPLANATORY NOTE**

Article II, Section 15 of the 1987 Constitution provides that the State shall protect and promote the right to health of the people and instill health consciousness among them.

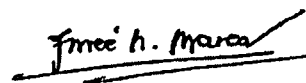
Article XIII, Sections 11 and 12 of the Constitution further mandate that "the State shall adopt an integrated and comprehensive approach to health development which shall endeavor to make essential goods, health and other social services available to all the people at affordable cost. There shall be priority for the needs of the under-privileged, sick, elderly, disabled, women, and children. The State shall endeavor to provide free medical care to paupers." "The State shall establish and maintain an effective food and drug regulatory system and undertake appropriate health manpower development and research, responsive to the country's health needs and problems."

The enactment of Republic Act No. 9502, otherwise known as the Universally Accessible Cheaper and Quality Medicines Act of 2008 has lowered the prices of medicines in the country. However, the Department of Health admitted that while the general trend in the prices of generic essential medicines have gone down in recent years, the Philippines is still paying higher prices when compared internationally. Generic drugs are still sold up to four times the international reference prices whereas branded innovator products are sold up to 22 times higher, especially in private hospitals and pharmacies.

This bill seeks to create an inter-agency Drug Price Regulatory Board (hereafter referred as the Board) to regulate the prices of drugs and medicines in the Philippines, in lieu of the Drugs and Medicines Price Monitoring and Regulation Authority of the Secretary of the Department of Health. The inter-agency composition of the Board ensures a broad spectrum of ideas, viewpoints and administrative powers in addressing the issue of affordability and accessibility of drugs and medicines in the country.

The bill also seeks to strengthen the procurement of cheaper drugs and medicines by the government through the creation of a trust fund to be utilized for parallel drug importation and other procurement arrangements. The Board is empowered to require pharmaceutical distributors to buy or obtain under any other form of arrangement, reasonable quantity of drugs and medicines procured by the government. The Board can also mandate up to 15% of a drug or medicine procurement of large pharmaceutical distributors to be allotted to a particular generic drug and medicine, thereby ensuring affordability and accessibility of medicines for our countrymen.

Thus, the passage of this bill is earnestly sought.

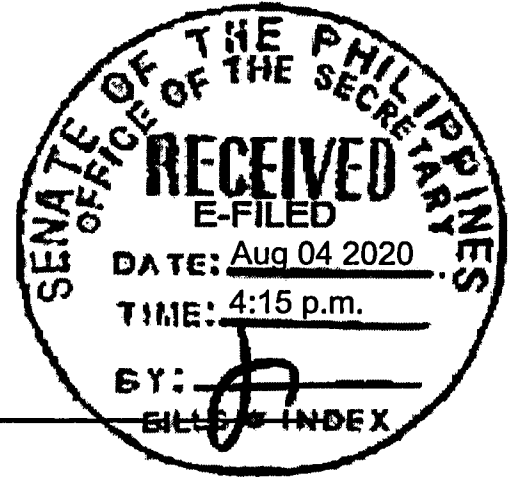
A handwritten signature in black ink that reads "Imee R. Marcos". The signature is written in a cursive style and is underlined with a single horizontal line.

**IMEE R. MARCOS**

EIGHTEENTH CONGRESS OF THE )  
REPUBLIC OF THE PHILIPPINES )  
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SENATE

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Introduced by **SEN. IMEE R. MARCOS**

**AN ACT ESTABLISHING THE DRUG PRICE REGULATORY BOARD TO  
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AMENDING FOR THE PURPOSE REPUBLIC ACT NO. 9502, OTHERWISE  
KNOWN AS THE "UNIVERSALLY ACCESSIBLE CHEAPER AND QUALITY  
MEDICINES ACT OF 2008" AND FOR OTHER PURPOSES**

*Be it enacted by the Senate and House of Representatives of the Philippines in Congress assembled:*

1           **SECTION 1.** Chapter 3 of Republic Act No. 9502 will now be titled, "*Drugs and*  
2 *Medicines Price Regulatory Board.*"  
3

4           **SEC. 2.** Section 17 of Republic Act No. 9502 is hereby deleted and a new  
5 Section 17 is hereby inserted to read as follows:  
6

7           **"SEC. 17. CREATION AND COMPOSITION OF THE DRUG**  
8 **PRICES REGULATION BOARD.**  
9

10          a) **THERE IS HEREBY CREATED THE DRUGS PRICES REGULATION**  
11 **BOARD, WHICH SHALL BE ATTACHED TO THE DEPARTMENT OF**  
12 **HEALTH AND COMPOSED OF SEVEN (7) MEMBERS AS FOLLOWS:**  
13

- 14           **1) SECRETARY OF HEALTH OR HIS DULY DESIGNATED**  
15 **REPRESENTATIVE AS CHAIRPERSON;**  
16           **2) SECRETARY OF TRADE AND INDUSTRY OR HIS DULY**  
17 **DESIGNATED REPRESENTATIVE AS VICE-CHAIRPERSON;**  
18           **3) DIRECTOR, FOOD AND DRUGS ADMINISTRATION OR HIS**  
19 **DULY DESIGNATED REPRESENTATIVE AS MEMBER;**

- 1           **4) CHAIRMAN, PHILIPPINE HEALTH INSURANCE**  
2           **CORPORATION AS MEMBER;**  
3           **5) ONE (1) ECONOMIST FROM THE ACADEME AS MEMBER; AND**  
4           **6) TWO (2) REPRESENTATIVES FROM THE CONSUMERS'**  
5           **SECTOR AS MEMBERS.**

- 6  
7           **b) THE MEMBERS OF THE BOARD REPRESENTING THE ACADEME**  
8           **AND THE CONSUMERS' SECTOR SHALL BE APPOINTED BY THE**  
9           **PRESIDENT OF THE PHILIPPINES AND SHALL SERVE FOR A**  
10           **TERM OF TWO (2) YEARS: *PROVIDED*, THAT THE**  
11           **REPRESENTATIVES FROM THE CONSUMERS' SECTOR SHALL NOT**  
12           **BE ELIGIBLE FOR REAPPOINTMENT FOR ANOTHER TERM."**

13  
14           **SEC. 3.** Section 18 of Republic Act No. 9502 is hereby deleted and a new  
15 Section 18 is hereby inserted to read as follows:

16  
17           **"SEC. 18. *POWERS OF THE BOARD.* – THE BOARD SHALL**  
18           **HAVE THE FOLLOWING POWERS:**

19  
20           **a)        *POWER TO DETERMINE THE MAXIMUM RETAIL PRICE***  
21           ***OF DRUGS OR MEDICINES SUBJECT TO PRICE REGULATION.* –**

22           **(1) UPON APPLICATION OR *MOTU PROPIO* WHEN THE PUBLIC**  
23           **INTEREST SO REQUIRES, THE BOARD SHALL HAVE THE**  
24           **POWER TO REGULATE THE RETAIL PRICES OF DRUGS AND**  
25           **MEDICINES LISTED UNDER SECTION 26 HEREOF,**  
26           **INCLUDING THEIR DOSAGE FORM AND PACKING, AND, IN**  
27           **ORDER THAT THEY SHALL BE MADE AVAILABLE TO THE**  
28           **PUBLIC AT AFFORDABLE RETAIL PRICE FROM THE**  
29           **DIFFERENT MANUFACTURERS, IMPORTERS, TRADERS,**  
30           **DISTRIBUTORS, WHOLESALERS OR RETAILERS AND AFTER**  
31           **A PROPER DETERMINATION AS THE BOARD MAY DEEM FIT,**  
32           **FIX FROM TIME TO TIME, BY PUBLICATION THE MAXIMUM**  
33           **RETAIL PRICE AT WHICH SUCH FORMULATIONS SHALL BE**  
34           **SOLD;**

35  
36           **(2) NO RETAILER SHALL SELL DRUGS AND MEDICINES AT A**  
37           **RETAIL PRICE EXCEEDING THE MAXIMUM RETAIL PRICE**  
38           **FIXED BY THE BOARD: *PROVIDED*, THAT UNTIL THE**  
39           **MAXIMUM RETAIL PRICE OF DRUGS AND MEDICINES**  
40           **SUBJECT TO PRICE REGULATION IS FIXED BY THE BOARD,**  
41           **THE RETAIL PRICE THEREOF SHALL BE THE PRICE WHICH**

1 PREVAILED IMMEDIATELY BEFORE THE EFFECTIVITY OF  
2 THIS ACT AND NO MANUFACTURER, IMPORTER, TRADER,  
3 DISTRIBUTOR, WHOLESALER OR RETAILER OF SUCH DRUG  
4 OR MEDICINE SHALL SELL THE SAME AT A RETAIL PRICE  
5 EXCEEDING THE PRICE PREVAILING IMMEDIATELY  
6 BEFORE THE EFFECTIVITY OF THIS ACT.  
7

8 FOR PURPOSES HEREOF, DRUGS AND MEDICINES  
9 SHALL INCLUDE BUT IS NOT LIMITED TO SINGLE- AND  
10 MULTI-INGREDIENT MEDICINES INCLUDED IN THE  
11 PHILIPPINE NATIONAL DRUG FORMULARY (PNDF)  
12 ESSENTIAL DRUG LIST AND SOLD UNDER THEIR GENERIC  
13 AND BRAND NAMES.  
14

15 **b) *POWER TO INCLUDE OTHER DRUGS OR MEDICINES IN THE LIST***  
16 ***SUBJECT TO PRICE REGULATION. – UPON APPLICATION OR***  
17 ***MOTU PROPIO WHEN THE PUBLIC INTEREST SO REQUIRES AND***  
18 ***AFTER PROPER DETERMINATION, THE BOARD MAY ORDER THE***  
19 ***INCLUSION OF DRUGS AND MEDICINES TO THE LIST SUBJECT***  
20 ***TO PRICE REGULATION UNDER SECTION 26 HEREOF.***  
21

22 **c) *POWER TO IMPLEMENT COST-CONTAINMENT AND OTHER***  
23 ***MEASURES. –***  
24

25 **1) THE BOARD SHALL HAVE THE POWER TO DETERMINE**  
26 **THE FAIR PRICE OF DRUGS OR MEDICINES FOR**  
27 **PURPOSES OF PUBLIC HEALTH INSURANCE AND**  
28 **GOVERNMENT PROCUREMENT; AND**

29 **2) THE BOARD SHALL HAVE THE POWER TO IMPLEMENT**  
30 **ANY OTHER MEASURES THAT THE GOVERNMENT MAY**  
31 **AVAIL OF TO EFFECTIVELY REDUCE THE COST OF DRUGS**  
32 **OR MEDICINES THAT SHALL INCLUDE, BUT NOT**  
33 **LIMITED TO, COMPETITIVE BIDDING, PRICE-VOLUME**  
34 **NEGOTIATIONS, PARALLEL DRUG IMPORTATION AND**  
35 **OTHER APPROPRIATE MECHANISMS THAT INFLUENCE**  
36 **SUPPLY, DEMAND, AND EXPENDITURES ON DRUGS AND**  
37 **MEDICINES.**

38 **3) THE BOARD SHALL HAVE THE POWER TO MANDATE UP**  
39 **TO 15% OF A DRUG OR MEDICINE PROCUREMENT OF**  
40 **LARGE PHARMACEUTICAL DISTRIBUTORS TO BE**  
41 **ALLOTTED TO A PARTICULAR GENERIC DRUG AND**

1 MEDICINE AND/OR REQUIRE PHARMACEUTICAL  
2 DISTRIBUTORS TO BUY OR OBTAIN UNDER ANY OTHER  
3 FORM OF ARRANGEMENTS, REASONABLE QUANTITY OF  
4 DRUGS AND MEDICINES PROCURED BY THE PHILIPPINE  
5 PHARMA PROCUREMENT, INC. SUCH DRUGS AND  
6 MEDICINES SHALL BE MADE AVAILABLE TO ALL  
7 BRANCHES OF THE SAID DISTRIBUTOR WHICH SHALL  
8 INFORM ANY BUYER OF THE AVAILABILITY, WITH  
9 CORRESPONDING PRICES, OF THESE DRUGS AND  
10 MEDICINES SO THAT THE BUYER MAY ADEQUATELY  
11 EXERCISE HIS/HER OPTION. THE LIST OF THESE DRUGS  
12 AND MEDICINES SHALL BE POSTED IN A CONSPICUOUS  
13 PLACE IN THE SAID BRANCHES.  
14

15 d) ***POWER TO IMPOSE ADMINISTRATIVE FINES AND PENALTIES.***

16 – AFTER DUE NOTICE AND HEARING, THE BOARD SHALL HAVE  
17 THE POWER TO SUSPEND OR REVOKE THE LICENSE TO  
18 OPERATE (LTO), PROFESSIONAL OR BUSINESS LICENSE, AS  
19 THE CASE MAY BE, OF ANY PERSON, MANUFACTURER,  
20 IMPORTER, TRADER, DISTRIBUTOR, WHOLESALER, RETAILER,  
21 OR ANY OTHER ENTITY, AND IMPOSE ADMINISTRATIVE FINES  
22 IN SUCH AMOUNT AS IT MAY DEEM REASONABLE WHICH  
23 SHALL IN NO CASE BE LESS THAN TWO HUNDRED THOUSAND  
24 PESOS (P 200,000.00) NOR MORE THAN FIVE MILLION PESOS  
25 (P 5,000,000.00) FOR VIOLATIONS OF THE MAXIMUM RETAIL  
26 PRICE FIXED PURSUANT TO THIS SECTION.  
27

28 e) ***OTHER POWERS NECESSARY TO IMPLEMENT PROVISIONS OF***

29 ***THIS CHAPTER-*** THE BOARD SHALL EXERCISE SUCH POWERS  
30 AND FUNCTIONS AS MAY BE NECESSARY TO IMPLEMENT AND  
31 ENFORCE THE PROVISIONS OF THIS CHAPTER OF THIS ACT,  
32 INCLUDING THE POWER TO REQUIRE THE PRODUCTION AND  
33 SUBMISSION OF RECORDS, DOCUMENTS, BOOKS OF  
34 ACCOUNT, BILLS OF LADING, INPUT DOCUMENTS, RECORDS  
35 OF PURCHASE AND SALE, FINANCIAL STATEMENTS, AND SUCH  
36 OTHER DOCUMENTS, INFORMATION AND PAPERS AS MAY BE  
37 NECESSARY TO ENABLE THE BOARD TO CARRY OUT ITS  
38 FUNCTIONS, DUTIES AND RESPONSIBILITIES. ACCORDINGLY,  
39 EVERY DECEMBER 31<sup>ST</sup> OF EVERY YEAR, EVERY  
40 MANUFACTURER, IMPORTER, TRADER, DISTRIBUTOR,  
41 WHOLESALER, AND RETAILER OF DRUG AND MEDICINE

1           **WHETHER INCLUDED IN OR EXCLUDED FROM THE LIST OF**  
2           **DRUGS AND MEDICINES THAT ARE SUBJECT TO PRICE**  
3           **REGULATION SHALL FURNISH THE BOARD A LIST,**  
4           **CONTAINING ON THE MINIMUM THE CORRESPONDING**  
5           **PRICES AND INVENTORY, OF ALL DRUGS AND MEDICINES IT**  
6           **MANUFACTURES, IMPORTS, TRADES, DISTRIBUTES,**  
7           **WHOLESALES, OR RETAILS AND ALL NECESSARY**  
8           **INFORMATION THAT THE BOARD MAY REQUIRE."**

9           **SEC. 4.** Section 19 of Republic Act No. 9502 is hereby deleted and a new  
10          Section 19 is hereby inserted to read as follows:

11          **"SEC. 19. MEETINGS OF THE BOARD — THE BOARD SHALL HOLD**  
12          **REGULAR MEETING EVERY QUARTER AND SUCH SPECIAL**  
13          **MEETINGS AS MAY BE NECESSARY UPON THE REQUEST OF THE**  
14          **CHAIRMAN OR UPON THE REQUEST OF AT LEAST TWO (2) OF ITS**  
15          **MEMBERS. THE BOARD MAY INVITE CONCERNED PUBLIC AND**  
16          **PRIVATE AGENCIES OR ENTITIES TO PARTICIPATE, COMPLEMENT,**  
17          **AND ASSIST IN THE PERFORMANCE OF ITS FUNCTIONS."**

18          **SEC. 5.** Section 20 of Republic Act No. 9502 is hereby deleted and a new  
19          Section 20 is hereby inserted to read as follows:

20          **"SEC. 20. CREATION OF A SECRETARIAT — THERE IS HEREBY**  
21          **CREATED A SECRETARIAT TO BE HEADED BY AN EXECUTIVE**  
22          **DIRECTOR TO SUPPORT THE BOARD IN CARRYING OUT ITS**  
23          **FUNCTIONS. THE BOARD SHALL PROVIDE FOR THE**  
24          **INSTITUTIONAL SETUP, QUALIFICATIONS, AND COMPENSATION**  
25          **OF THE EMPLOYEES COMPOSING THE SECRETARIAT IN**  
26          **ACCORDANCE WITH EXISTING CIVIL SERVICE AND CAREER**  
27          **EXECUTIVE SERVICE RULES AND REGULATIONS AND CONSISTENT**  
28          **WITH THE PROVISION OF THE SALARY STANDARDIZATION LAW**  
29          **FOR GOVERNMENT PERSONNEL, AND DETERMINE THE SIZE AND**  
30          **COMPOSITION OF THE SECRETARIAT."**

31          **SEC. 6.** Section 21 of Republic Act No. 9502 is hereby deleted and a new  
32          Section 21 is hereby inserted to read as follows:

33          **"SEC. 21. PROCEDURES FOR INQUIRIES, STUDIES, HEARINGS,**  
34          **INVESTIGATIONS, AND PROCEEDINGS. —ALL INQUIRIES,**  
35          **STUDIES, HEARINGS, INVESTIGATIONS AND PROCEEDINGS**  
36          **CONDUCTED BY THE BOARD SHALL BE GOVERNED BY THE RULES**

1           **ADOPTED BY THE BOARD, AND IN THE CONDUCT THEREOF SHALL**  
2           **NOT BE BOUND BY THE TECHNICAL RULES OF EVIDENCE."**

3           **SEC. 7.** Section 22 of Republic Act No. 9502 is hereby deleted and a new  
4 Section 22 is hereby inserted to read as follows:

5           **"SEC. 22. EFFECTIVITY AND REVIEW OF THE DECISIONS OR**  
6           **ORDERS OF THE BOARD. — ALL DECISIONS OR ORDERS OF THE**  
7           **BOARD PURSUANT TO SECTION 18 HEREOF, SHALL BE**  
8           **IMMEDIATELY OPERATIVE.**

9           **A PARTY ADVERSELY AFFECTED BY A DECISION, ORDER OR**  
10          **RULING OF THE BOARD MAY, WITHIN THIRTY (30) DAYS FROM**  
11          **NOTICE OF SUCH DECISION, ORDER OR RULING, OR IN CASE OF A**  
12          **DENIAL OF A MOTION FOR RECONSIDERATION THEREOF, WITHIN**  
13          **FIFTEEN (15) DAYS AFTER NOTICE OF SUCH DENIAL, FILE AN**  
14          **APPEAL WITH THE COURT OF APPEALS, WHICH SHALL HAVE**  
15          **JURISDICTION TO REVIEW SUCH DECISION, ORDER OR RULING.**

16          **THE FILING OF A PETITION FOR A WRIT OF CERTIORARI OR**  
17          **OTHER SPECIAL REMEDIES IN THE SUPREME COURT SHALL IN NO**  
18          **CASE SUPERSEDE OR STAY ANY DECISION, ORDER OR RULING OF**  
19          **THE BOARD, UNLESS THE SUPREME COURT SHALL SO DIRECT, AND**  
20          **THE PETITIONER MAY BE REQUIRED BY THE SUPREME COURT TO**  
21          **GIVE BOND IN SUCH FORM AND OF SUCH AMOUNT AS MAY BE**  
22          **DEEMED PROPER."**

23  
24          **SEC. 8.** Section 26 of Republic Act No. 9502 is hereby amended to read as  
25 follows:

26  
27          **"SECTION 26. *Display of Maximum Retail Price Fixed [and***  
28          ***approved by order of the President of the Philippines] by the BOARD***  
29          ***for Drugs or Medicines Subject to Price Regulation.* — (a) Within a**  
30          **reasonable period as may be determined by the [Secretary of the Department**  
31          **of Health] BOARD, and: *Provided,* That it conforms to existing drug product**  
32          **labeling requirements, every manufacturer, importer, distributor, wholesaler,**  
33          **trader, or retailer of a drug and medicine intended for sale shall display the**  
34          **retail price which shall not exceed the maximum retail price fixed by the**  
35          **Board. The maximum retail price shall be printed on the label of the**  
36          **immediate container of the drug and medicine and the minimum pack thereof**  
37          **offered for retail sale with the words "RETAIL PRICE NOT TO EXCEED"**  
38          **preceding it, and "UNDER DRUG PRICE REGULATION" on a red strip**  
39          **PROVIDED THAT IN CASE OF A CONTAINER CONSISTING OF**



1 **SMALLER SALEABLE PACKS, THE RETAIL PRICE OF SUCH SMALLER**  
2 **PACK SHALL ALSO BE DISPLAYED ON THE LABEL OF EACH SMALLER**  
3 **PACK AND SUCH PRICE SHALL NOT BE MORE THAN THE *PRO RATA***  
4 **RETAIL PRICE OF THE MAIN PACK ROUNDED OFF TO THE NEAREST**  
5 **CENTAVO.**

6  
7 (b) Within a period as may be determined by the [Secretary of the  
8 Department of Health] **BOARD** from time to time, every manufacturer,  
9 importer, or trader shall issue a price list to wholesalers, distributors, retailers  
10 and to the [Secretary of the Department of Health] **BOARD**, indicating the  
11 retail price, the maximum retail price, and such other information as may be  
12 required by the [Secretary of the Department of Health] **BOARD**.

13  
14 **SEC. 9.** A new Section 26-A is hereby inserted to read as follows:

15  
16 **"SECTION 26-A. *Display of Price and Price List of Drugs or Medicines***  
17 ***Excluded from the List Subject to Price Regulation.*** – Every  
18 manufacturer, importer, trader, distributor, wholesaler, or retailer of a drug  
19 or medicine excluded from the list subject to price regulation under Section  
20 23 hereof shall display in indelible print mark on the label of the immediate  
21 container of the drug or medicine and the minimum pack thereof offered for  
22 retail sale, the words "**NOT UNDER PRICE REGULATION**" on green strip.

23  
24 **SEC. 10.** Section 28 of R.A 9502 is hereby deleted and a new Section 28 is  
25 hereby inserted to read as follows:

26  
27 **"Sec. 28. *Creation of a Trust Fund*** A trust fund in the amount of  
28 Five Hundred Million Pesos (P 500,000,000.00) is hereby established to be  
29 administered by the Board. The fund shall serve as revolving fund to be  
30 utilized for parallel drug importation and other procurement arrangements to  
31 lower the cost of drugs and medicines through the Philippine Pharma  
32 Procurement, Inc."

33  
34 **SEC. 11.** Section 30 of R.A. 9502 is hereby amended to read as  
35 follows:

36  
37 **"SEC. 30. Reportorial and Public Notice Requirements.** — (a) The  
38 [Secretary of the Department of Health] **BOARD** shall submit a bi-annual  
39 Monitoring Report of its performance on the implementation of this Act to  
40 the Office of the President. This report submitted to the Office of the

1 President shall be published in a newspaper of general circulation within thirty  
2 (30) days upon submission.

3  
4 X x x x

5  
6 (c) The order of the [President of the Philippines] **BOARD** imposing  
7 maximum retail prices on drugs and medicines, including the conditions  
8 implementing it, shall be published within fifteen (15) days from issuance in  
9 at least two (2) newspapers of general circulation. All wholesalers,  
10 manufacturers, distributors, importers, or traders shall have a copy of the  
11 order of the [President of the Philippines] **BOARD** and provide the same to  
12 their clients and customers for every transaction.

13  
14 X x x x"

15  
16 **SEC. 12. Appropriations.** – The amount of Five Hundred Million Pesos (P  
17 500,000,000.00) as a trust fund under Section 28 hereof and the amount necessary  
18 to carry out the functions of the Board shall be included in the annual General  
19 Appropriations Act.

20  
21 **SEC. 13. Separability Clause.** – Should any provision herein be declared  
22 unconstitutional, the same shall not affect the validity of other provisions of this Act.

23  
24 **SEC. 14. Repealing Clause.** – All laws, decrees, orders, rules, and regulations  
25 or other issuances of parts inconsistent with the provisions of this Act are hereby  
26 repealed or modified accordingly.

27  
28 **SEC. 15. Effectivity Clause.** – This Act shall take effect fifteen (15) days  
29 after its publication in the *Official Gazette* or in any two (2) newspapers of general  
30 circulation in the Philippines.

*Approved,*