



The President
Palikir, Pohnpei
Federated States of Micronesia



June 21, 2022

The Honorable Wesley W. Simina
Speaker
Twenty-Second Congress
Federated States of Micronesia
Palikir, Pohnpei FM 96941

PRESIDENTIAL COMM. NO. 22-288
FSM CONGRESS

Dear Speaker Simina:

I am pleased to transmit the following act, which I have signed to become Public Law No. 22-136 :

Congressional Act No. 22-112, entitled: "AN ACT TO FURTHER AMEND TITLE 41 OF THE CODE OF THE FEDERATED STATES OF MICRONESIA (ANNOTATED), AS AMENDED, BY CREATING A NEW CHAPTER 13 TO ESTABLISH THE FSM SAFE PHARMACEUTICAL ACT OF 2022, ESTABLISH THE PHARMACEUTICAL UNIT UNDER THE DEPARTMENT OF HEALTH, ADOPT CRITERIA FOR THE FSM APPROVED MEDICINES LIST AND COMPETENT JURISDICTIONS DESIGNATION, ESTABLISH THE PHARMACEUTICAL UNIT LICENSURE AND PHARMACEUTICAL PRODUCT REGISTRATION PROCESS, AND AUTHORIZE THE SECRETARY OF HEALTH TO SUSPEND OR REVOKE ANY PHARMACEUTICAL UNIT LICENSE OR PRODUCT REGISTRATION APPROVAL FOR CAUSE, AND FOR OTHER PURPOSES."

I thank the 22nd FSM Congress for the passage of this act.

Sincerely,

David W. Panuelo
President

Xc: Chief Justice, FSM Supreme Court



Office of the Chief Clerk

CONGRESS OF THE FEDERATED STATES OF MICRONESIA

P.O. Box PS 3

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PRESIDENTIAL COMM. NO. 22-288
FSM CONGRESS

May 21, 2022

His Excellency David W. Panuelo
President
Federated States of Micronesia
Palikir, Pohnpei FM 96941

Dear President Panuelo:

I have the honor to transmit herewith Congressional Act No. 22-112, "AN ACT TO FURTHER AMEND TITLE 41 OF THE CODE OF THE FEDERATED STATES OF MICRONESIA (ANNOTATED), AS AMENDED, BY CREATING A NEW CHAPTER 13 TO ESTABLISH THE FSM SAFE PHARMACEUTICAL ACT OF 2022, ESTABLISH THE PHARMACEUTICAL UNIT UNDER THE DEPARTMENT OF HEALTH, ADOPT CRITERIA FOR THE FSM APPROVED MEDICINES LIST AND COMPETENT JURISDICTIONS DESIGNATION, ESTABLISH THE PHARMACEUTICAL UNIT LICENSURE AND PHARMACEUTICAL PRODUCT REGISTRATION PROCESS, AND AUTHORIZE THE SECRETARY OF HEALTH TO SUSPEND OR REVOKE ANY PHARMACEUTICAL UNIT LICENSE OR PRODUCT REGISTRATION APPROVAL FOR CAUSE, AND FOR OTHER PURPOSES", which was passed by the Twenty-Second Congress of the Federated States of Micronesia, Fourth Regular Session, 2022, by a two-thirds vote of all the State delegations as required and as duly certified.

Sincerely yours,

A handwritten signature in black ink, appearing to read "Jessicalynn Reyes", is written over a horizontal line.

Jessicalynn Reyes
Chief Clerk, Congress of the
Federated States of Micronesia

Enclosures

RECEIVED
MAY 27 2022
Office of the
President FSM



PRESIDENTIAL COMM. NO. 22-288
FSM CONGRESS

TWENTY-SECOND CONGRESS OF THE
FEDERATED STATES OF MICRONESIA
FOURTH REGULAR SESSION
MAY 04 – [23] 27, 2022
(EXTENDED BY C.R. NO. 22-130, ADOPTED 05/23/22)

An Act

TO FURTHER AMEND TITLE 41 OF THE CODE OF THE FEDERATED STATES OF MICRONESIA (ANNOTATED), AS AMENDED, BY CREATING A NEW CHAPTER 13 TO ESTABLISH THE FSM SAFE PHARMACEUTICAL ACT OF 2022, ESTABLISH THE PHARMACEUTICAL UNIT UNDER THE DEPARTMENT OF HEALTH, ADOPT CRITERIA FOR THE FSM APPROVED MEDICINES LIST AND COMPETENT JURISDICTIONS DESIGNATION, ESTABLISH THE PHARMACEUTICAL UNIT LICENSURE AND PHARMACEUTICAL PRODUCT REGISTRATION PROCESS, AND AUTHORIZE THE SECRETARY OF HEALTH TO SUSPEND OR REVOKE ANY PHARMACEUTICAL UNIT LICENSE OR PRODUCT REGISTRATION APPROVAL FOR CAUSE, AND FOR OTHER PURPOSES.

INTRODUCED BY SENATOR: FERNY S. PERMAN

DATE: MAY 04, 2022

REFERRED TO: COMMITTEE ON WAYS AND MEANS

S.C.R. NO. 22-33 – MAY 19, 2022

FIRST READING: – MAY 23, 2022

SECOND READING: – MAY 24, 2022

A handwritten signature in black ink, appearing to read "Jessicalynn Reyes", is written over a horizontal line.

Jessicalynn Reyes
Chief Clerk, FSM Congress



Office of the Speaker

CONGRESS OF THE FEDERATED STATES OF MICRONESIA

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PRESIDENTIAL COMM. NO. 22-288
FSM CONGRESS

ACT NO. 22-112

(CONGRESSIONAL BILL NO. 22-166, C.D.1)

We hereby certify that on May 24 the foregoing act passed
Second and Final Reading of the Twenty-Second Congress of the
Federated States of Micronesia, Fourth Regular Session, 2022,
by a two-thirds vote of all the State delegations as required
under article IX, section 20, of the Constitution of the
Federated States of Micronesia.

A handwritten signature in black ink, appearing to read "Wesley W. Simina".

Wesley W. Simina
Speaker
Congress of the
Federated States of Micronesia

A handwritten signature in black ink, appearing to read "Jessicalynn Reyes".

Jessicalynn Reyes
Chief Clerk
Congress of the
Federated States of Micronesia

TWENTY-SECOND CONGRESS OF THE FEDERATED STATES OF MICRONESIA

FOURTH REGULAR SESSION, 2022

CONGRESSIONAL BILL NO. 22-166 C.D.1

AN ACT

To further amend title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, by creating a new chapter 13 to establish the FSM Safe Pharmaceutical Act of 2022, establish the Pharmaceutical Unit under the Department of Health, adopt criteria for the FSM Approved Medicines List and competent jurisdictions designation, establish the Pharmaceutical Unit licensure and pharmaceutical product registration process, and authorize the Secretary of Health to suspend or revoke any Pharmaceutical Unit license or product registration approval for cause, and for other purposes.

BE IT ENACTED BY THE CONGRESS OF THE FEDERATED STATES OF MICRONESIA:

1 Section 1. Title 41 of the Code of the Federated States of
2 Micronesian (Annotated), as amended, is hereby amended by creating
3 a new chapter 13 entitled: "FSM Safe Pharmaceutical Act of 2022".

4 Section 2. Chapter 13 of title 41 of the Code of the
5 Federated States of Micronesia (Annotated), as amended, is hereby
6 amended by inserting a new subchapter 1 entitled: "General
7 Provisions".

8 Section 3. Chapter 13 of title 41 of the Code of the
9 Federated States of Micronesia (Annotated), as amended, is hereby
10 amended by inserting a new section 1301 of subchapter 1 to read as
11 follows:

12 "Section 1301. Short title. This Act may be referred to
13 as the Safe Pharmaceutical Act."

14 Section 4. Chapter 13 of title 41 of the Code of the
15 Federated States of Micronesia (Annotated), as amended, is hereby

1 amended by inserting a new section 1302 of subchapter 1 to read as
2 follows:

3 "Section 1302. Statement of Policy. It is hereby
4 declared as a policy of the Federated States of
5 Micronesia:

6 1. That all people have the right to access quality,
7 safe and effective medicines;

8 2. The establishment and enforcement of import
9 controls on all pharmaceuticals is necessary to ensure
10 acceptable standards of quality, safety and efficacy of
11 pharmaceuticals entering the country; and ensure the
12 practices of all persons, businesses, entities and
13 establishments involved in the importation of
14 pharmaceuticals into the FSM comply with the acceptable
15 standards of quality, safety and efficacy; and

16 3. The national government and appropriate
17 government departments shall, to the extent possible,
18 cooperate with regulatory authorities in other countries
19 as appropriate, to strengthen pharmaceutical import
20 controls and align regulatory processes where needed to
21 tackle public health emergencies, and address the
22 proliferation of substandard, falsified and unlicensed
23 pharmaceuticals entering the FSM."

24 Section 5. Chapter 13 of title 41 of the Code of the
25 Federated States of Micronesia (Annotated), as amended, is hereby

1 amended by inserting a new section 1303 of subchapter 1 to read as
2 follows:

3 "Section 1303. Definitions: For the purposes of this
4 title, the following terms shall be given the meanings
5 described herein:

6 (1) "Active Pharmaceutical Ingredient" (API) is the
7 chemical substance contained in a pharmaceutical, which
8 is responsible for its therapeutic effect. Some
9 pharmaceuticals contain more than one active ingredient
10 (combination product).

11 (2) "Authorized port of entry" means a port of entry
12 designated by the Secretary of Justice under Section 202
13 of Title 18 of the Code of the FSM.

14 (3) "Certificate of pharmaceutical product (CPP)"
15 means a certificate issued by the authorized body of the
16 exporting country that satisfies the pharmaceutical
17 verification format standards to permit importation into
18 the FSM as determined by the Secretary of Health.

19 (4) "Competent jurisdictions" means countries with
20 stringent and operational regulatory system where
21 medicines can be imported into the FSM as determined by
22 the Secretary of Health.

23 (5) "Customs Administration" means the Customs and
24 Tax Administration under the FSM Department of Finance
25 and Administration.

(6) "Department of Health" means the Department of Health and Social Affairs.

(7) "Distribution" means the division and movement of pharmaceuticals from the port of entry to the Establishment or end user thereof, by means of various transport methods or storage.

(8) "Distributor" means an individual, company or legal entity distributing or seeking to distribute a pharmaceutical.

(9) "Donation" means the act by which organizations, institutions, international development partners, non-government organizations and other legal entities provide pharmaceuticals to the FSM for free and for specific use, such as in the case of emergency or for humanitarian purposes.

(10) "Establishment" means an entity in the FSM that engages in the importation of pharmaceuticals and/or active pharmaceutical ingredients into the FSM, including but not limited to:

- a. Wholesalers;
- b. Distributors;
- c. Pharmacies;
- d. Importers;
- e. Exporters;
- f. Manufacturers; and

1 g. Warehouse operators.

2 (11) "Exportation" means the lawful process of
3 sending medicines out of the FSM by, sea or air.

4 (12) "Exporter" means an individual, company or legal
5 entity that exports pharmaceuticals.

6 (13) "FSM Approved Medicines List" means a list of
7 pharmaceuticals determined by the Secretary of Health to
8 meet the needs of the FSM population and satisfy the
9 pharmaceutical product registration approval criteria for
10 importation into the FSM.

11 (14) "Importation" means the lawful process of
12 bringing medicines into the FSM, by sea or air.

13 (15) "Importer" means an individual, company or
14 similar legal entity importing or seeking to import
15 pharmaceuticals.

16 (16) "Inspectoral" means an official examination,
17 usually conducted on-site by the relevant authority to
18 determine compliance to regulations, standards and
19 practices by Establishments, and/or any other entity
20 engaged in the import of pharmaceuticals into the FSM.

21 (17) "Manufacturing" means all operations of
22 procuring supply, production, packaging, repackaging,
23 labeling, relabeling, quality control, release, storage
24 and distribution of active pharmaceutical ingredients and
25 related controls.

(18) "Over-the-counter medicines (non-prescription medicines)" means medicines sold from licensed dealers without professional supervision and prescription that are suitable for self-medication for minor disease and symptoms.

(19) "Pharmaceutical" means any substance or medical product for human or veterinary use that is intended to modify or explore physiological systems or pathological states for the benefit of the recipient. The term "pharmaceutical" includes any pharmaceutical product, drug, medicine, vaccine, biopharmaceuticals, blood and blood products, active pharmaceutical ingredient, and any other products with therapeutic effect.

(20) "Prescription" means an order mostly in written form by a licensed health care professional to a pharmacist or other therapist for a pharmaceutical or medicine to be provided to the health care professional's patient.

(21) "Procurement" means the process of acquiring pharmaceuticals, including those obtained by purchase and/or donation.

(22) "Quality assurance" means the comprehensive review of the pharmaceutical supply system and process based on scientifically accepted standards in the industry to assess the quality of the pharmaceutical.

(23) "Sampling" means an operations designed to obtain a representative portion of a pharmaceutical product, based on an appropriate statistical procedure, for a defined purpose.

(24) "Secretary of Health" means the Secretary of Health and Social Affairs.

(25) "Wholesale" means all activities consisting of procuring, holding, or supplying pharmaceuticals for import or export.

(26) "Wholesaler" means an individual, company or similar legal entity engaged in the wholesale of pharmaceuticals."

Section 6. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by creating a new subchapter 2 entitled: "Scope of the Law".

Section 7. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by inserting a new section 1304 of subchapter 2 to read as follows:

"Section 1304. Scope of Law.

(1) Pharmaceutical Products.

All pharmaceuticals imported into the FSM shall be regulated under this Act. Any drug, medicine, or health supplement imported into the FSM with a therapeutic claim

that is not scientifically verifiable shall be treated and regulated as a pharmaceutical under this Act.

(2) Pharmaceutical Activities.

All Establishment pharmaceutical activities related to the importation of pharmaceuticals into the FSM shall be regulated under this Act. Only Establishments licensed by the Pharmaceutical Unit are eligible to import pharmaceuticals at authorized ports of entry in compliance with any Pharmaceutical Unit licensure, pharmaceutical product registration and approval process, and procurement, storage, record-keeping and disposal requirements under Section 1305.

(3) Exempt Pharmaceuticals and Activities.

(a) The regulation of pharmaceuticals and activities under this Act does not apply to natural or indigenous medicines native to the FSM.

(b) The regulation of pharmaceuticals and pharmaceutical activities under this Act does not apply to the importation of pharmaceuticals into the FSM for personal use subject to the following requirements:

(i) Pharmaceutical is an over-the-counter medicine that is not for the treatment of a serious medical condition and there is no known significant health risk; or

(ii) Pharmaceutical is prescribed by a

licensed doctor under the following conditions:

(A) pharmaceutical is accompanied by a prescription from a FSM licensed doctor or prescription from a foreign country licensed doctor with certification that the pharmaceutical is a continuation of medical treatment performed by the foreign licensed doctor in the same foreign country where the doctor is licensed;

(B) the consumer of the prescribed pharmaceutical affirms in writing that the pharmaceutical is for personal use and will not be commercialized or distributed to other persons in the FSM; and

(C) the quantity on the pharmaceutical Prescription is not more than a three-month supply.

(4) Establishment Requirements.

Establishments shall be licensed by the Pharmaceutical Unit in order to be eligible to import pharmaceuticals into the FSM. Establishments shall register any pharmaceutical it intends to import with the Pharmaceutical Unit and receive pharmaceutical product registration approval from the Pharmaceutical Unit before importing the pharmaceutical into the FSM."

Section 8. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by creating a new subchapter 3 entitled: "Administration."

Section 9. Chapter 13 of title 41 of the Code of the

1 Federated States of Micronesia (Annotated), as amended, is hereby
2 amended by inserting a new section 1305 of subchapter 3 to read as
3 follows:

4 "Section 1305. Pharmaceutical Unit.

5 (1) The Secretary of Health shall establish the
6 Pharmaceutical Unit under the Department of Health to be
7 headed by a coordinator, otherwise known as the
8 Pharmaceutical Unit Coordinator, within 90 business days
9 of enactment of this Act.

10 (2) The Pharmaceutical Unit shall have
11 administrative, regulatory, inspectoral, and quality
12 assurance functions.

13 (3) Within 90 business days of establishment of the
14 Pharmaceutical Unit, the Pharmaceutical Unit shall adopt
15 criteria for the FSM Approved Medicines List and
16 standards for the competent jurisdiction designation, and
17 submit to the Secretary of Health for approval. The
18 Secretary of Health shall review and make a decision on
19 the FSM Approved Medicines List criteria and competent
20 jurisdiction designation standards proposed by the
21 Pharmaceutical Unit within 30 business days. The
22 Secretary of Health must approve the FSM Approved
23 Medicines List criteria and competent jurisdiction
24 designation standards prior to regulation.

25 (4) Within 90 business days of establishment of the

1 Pharmaceutical Unit, the Pharmaceutical Unit shall
2 develop the Establishment licensure and pharmaceutical
3 product registration process, and submit to the Secretary
4 Health for approval. The Secretary of Health shall
5 review and make a decision on the Establishment licensure
6 and pharmaceutical product registration process proposed
7 by the Pharmaceutical Unit within 30 business days. The
8 Secretary of Health must approve the Establishment
9 licensure and pharmaceutical product registration process
10 prior to regulation.

11 (5) Upon compliance with subsection 4 of this
12 Section, the Pharmaceutical Unit shall have the authority
13 to implement the Establishment Licensure, and
14 pharmaceutical product registration process requirements
15 and conditions under Section 1309 and Section 1310.

16 (6) The Pharmaceutical Unit Coordinator may call
17 upon independent experts and/or technical partners to
18 assist the Pharmaceutical Unit in the development of
19 criteria for the FSM Approved Medicines List and
20 competent jurisdiction designations, Establishment
21 licensure and pharmaceutical product registration
22 process, and evaluation of pharmaceuticals for product
23 registration purposes."

24 Section 10. Chapter 13 of title 41 of the Code of the
25 Federated States of Micronesia (Annotated), as amended, is hereby

1 amended by creating a new subchapter 4 entitled: "Regulation
2 Authority."

3 Section 11. Chapter 13 of title 41 of the Code of the
4 Federated States of Micronesia (Annotated), as amended, is hereby
5 amended by inserting a new section 1306 of subchapter 4 to read as
6 follows:

7 "Section 1306. Regulation through Rule-Making.

8 The Secretary of Health shall implement this Act by
9 regulation in accordance with administrative rule-making
10 procedures under Chapter 1 of Title 17 of the Code of the
11 FSM."

12 Section 12. Chapter 13 of title 41 of the Code of the
13 Federated States of Micronesia (Annotated), as amended, is hereby
14 amended by inserting a new section 1307 of subchapter 4 to read as
15 follows:

16 "Section 1307. Certificate of Pharmaceutical Product
17 Requirements.

18 The Secretary of Health shall establish the Certificate
19 for Pharmaceutical Product (CPP) form requirements in the
20 World Health Organization recommended format or the
21 equivalent, and implement regulations accordingly."

22 Section 13. Chapter 13 of title 41 of the Code of the
23 Federated States of Micronesia (Annotated), as amended, is hereby
24 amended by inserting a new section 1308 of subchapter 4 to read as
25 follows:

1 "Section 1308. FSM Approved Medicines List.

2 (1) The Secretary of Health shall establish the FSM
3 Approved Medicines List and determine the medicines on the
4 FSM Approved Medicines List. The Secretary of Health
5 shall consider the Pharmaceutical Unit recommendations for
6 pharmaceuticals to add, remove from, or modify on the FSM
7 Approved Medicines List.

8 (2) The Secretary of Health shall review the FSM
9 Approved Medicines List and the designation of competent
10 jurisdictions every five years or upon the Secretary of
11 Health certification to Congress that imminent peril to
12 the public health, safety, or welfare requires immediate
13 review and changes to the Approved Medicines List and/or
14 competent jurisdiction designations. Upon review of the
15 FSM Approved Medicines List and the competent jurisdiction
16 designations, the Secretary of Health shall determine
17 whether pharmaceuticals and jurisdictions will be added,
18 removed, or modified, respectively.

19 (3) Only pharmaceuticals listed on the FSM Approved
20 Medicines List from competent jurisdictions can be
21 imported into the FSM by licensed Establishments without
22 the specific pharmaceutical registration approval from
23 the Pharmaceutical Unit.

24 (4) The Secretary of Health may add pharmaceuticals
25 to the FSM Approved Medicines List upon certification of

need by the States government health authorities and
review of the pharmaceutical."

Section 14. Chapter 13 of title 41 of the Code of the
Federated States of Micronesia (Annotated), as amended, is hereby
amended by inserting a new section 1309 of subchapter 4 to read as
follows:

"Section 1309. Establishment licensing requirements.

(1) The Secretary of Health or his designee shall have
the authority to regulate the licensure requirements for
Establishments.

(2) All Establishments shall be licensed by the
Pharmaceutical Unit in order to be eligible to import
pharmaceuticals into the FSM. Establishments are
prohibited from importing pharmaceuticals without a valid
license from the Pharmaceutical Unit.

(3) All Establishments shall comply with the licensure
standards and conditions set by the Secretary of Health
or his designee including but not limited to unannounced
random collection of a sample of the pharmaceutical at
the authorized ports of entry for quality assurance and
testing purposes.

(4) The Secretary of Health, with input from the
Pharmaceutical Unit Coordinator, shall determine the fee
and rules for Establishments to apply for licensure from
the Pharmaceutical Unit."

1 Section 15. Chapter 13 of title 41 of the Code of the
2 Federated States of Micronesia (Annotated), as amended, as
3 amended, is hereby amended by inserting a new section 1310 of
4 subchapter 4 to read as follows:

5 "Section 1310. Pharmaceutical product registration
6 system.

7 (1) The Secretary of Health or his designee shall
8 have the authority to regulate the pharmaceutical product
9 registration requirements. Pharmaceuticals registered in
10 competent jurisdictions as determined by the Secretary of
11 Health and designated on the FSM Approved Medicines List
12 may be exempt from the pharmaceutical registration review
13 requirements.

14 (2) The Secretary of Health, with input from the
15 Pharmaceutical Unit Coordinator, shall establish the
16 criteria and conditions for pharmaceutical product
17 registration. The criteria and conditions for
18 pharmaceutical product registration shall include but are
19 not limited to:

20 (a) only licensed Establishments eligible to
21 apply for pharmaceutical product registration with the
22 Pharmaceutical Unit;

23 (b) procurement, storage and disposal
24 requirements for pharmaceuticals; and

25 (c) record-keeping requirements for

pharmaceuticals.

(3) The Secretary of Health or his designee shall regulate the criteria and procedure for registration of new chemical compounds and/or variations to pharmaceuticals currently registered in the pharmaceutical product registration system.

(4) The Secretary of Health, with input from the Pharmaceutical Unit Coordinator, shall determine the fee and rules for Establishments to apply for pharmaceutical registration with the Pharmaceutical Unit."

Section 16. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by inserting a new section 1311 of subchapter 4 to read as follows:

"Section 1311. Enforcement and Penalties.

(1) Suspension and Revocation of Establishment License.

The Secretary of Health shall have the authority to suspend or revoke for cause an Establishment license.

The Establishment shall have the right to request review and/or administrative hearing on the license suspension or revocation in accordance with Chapter 1 of Title 17 of the Code of the FSM.

(2) Suspension and Revocation of Approved Pharmaceutical Product Registration.

1 The Secretary of Health shall have the authority to
2 suspend and/or revoke for cause any approved
3 pharmaceutical product registration. The Establishment
4 shall have the right to request review and/or
5 administrative hearing on the product registration
6 suspension or revocation in accordance with Chapter 1 of
7 Title 17 of the Code of the FSM."

8 Section 17. Chapter 13 of title 41 of the Code of the
9 Federated States of Micronesia (Annotated), as amended, is hereby
10 amended by inserting a new section 1312 of subchapter 4 to read as
11 follows:

12 "Section 1312. Entry of Pharmaceuticals for Public
13 Health Emergency and Life Saving Assistance.

14 (1) The Secretary of Health shall establish and
15 facilitate a streamlined process with the Assistant
16 Secretary for Customs to permit licensed Establishments
17 to import pharmaceuticals not on the FSM Approved
18 Medicines List, but from competent jurisdictions, for
19 public health and life-saving emergencies

20 (2) The Secretary of Health shall only permit licensed
21 Establishments to import pharmaceuticals not on the FSM
22 Approved Medicines List, but from competent
23 jurisdictions, upon written certification to Congress
24 that life-saving assistance or imminent peril to the
25 public health, safety, or welfare requires the immediate

entry of the pharmaceutical outside of the processes under Section 1309 of this Act.

(3) The Pharmaceutical Unit shall conduct a review of any pharmaceutical imported under this Section in accordance with the established pharmaceutical registration criteria within 30 calendar days of the pharmaceutical importation into the FSM. If the pharmaceutical does not satisfy the pharmaceutical registration criteria, the Secretary of Health shall recall the pharmaceutical."

Section 18. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by inserting a new section 1313 of subchapter 4 to read as follows:

"Section 1313: Donations.

The Secretary of Health shall only accept donations of pharmaceuticals on the FSM Approved Medicines List from competent jurisdictions and pharmaceuticals that will not expire for at least 1 year."

Section 19. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by creating a new subchapter 5 entitled: "Confidentiality and Whistleblower Protections".

Section 20. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby

1 amended by inserting a new section 1314 of subchapter 5 to read as
2 follows:

3 "Section 1314. Confidentiality of Records and
4 Whistleblower Protections.

5 (1) The Department of Health shall keep confidential
6 all information from any source on pharmaceutical
7 activities regulated under this Act, except in response
8 to a FSM Department administrative order, FSM subpoena or
9 court order, request from Congress pursuant to its
10 oversight powers, or request from the information source
11 for access to their own records in accordance with policy
12 and procedures established by regulations and
13 legislation.

14 (2) The Secretary of Health shall establish
15 whistleblower protections under this Act by regulation,
16 policy, and/or procedure."

17 Section 21. Chapter 13 of title 41 of the Code of the
18 Federated States of Micronesia (Annotated), as amended, is hereby
19 amended by creating a new subchapter 6 entitled: "Prohibited
20 Pharmaceutical Activities."

21 Section 22. Chapter 13 of title 41 of the Code of the
22 Federated States of Micronesia (Annotated), as amended, is hereby
23 amended by inserting a new section 1315 of subchapter 6 to read as
24 follows:

25 "Section 1315. Prohibited Pharmaceutical Activities.

(1) Establishments are prohibited from acting as an internet pharmacy for the importation of pharmaceuticals outside the Pharmaceutical Unit product registration and review process.

(2) Establishments are prohibited from using the personal use exemption under Section 1304(3) for the importation of pharmaceuticals outside the Pharmaceutical Unit product registration process under Section 1310 of this Act.

(3) Establishments licensed by the Pharmaceutical Unit are prohibited from manufacturing pharmaceuticals in the FSM and importing pharmaceuticals and/or active pharmaceutical ingredients to manufacturer pharmaceuticals in the FSM."

Section 23. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by creating a new subchapter 7 entitled: "Civil and Criminal Actions".

Section 24. Chapter 13 of title 41 of the Code of the Federated States of Micronesia (Annotated), as amended, is hereby amended by inserting a new section 1316 of subchapter 7 to read as follows:

"Section 1316. Civil and Criminal Actions.

This Act shall not be construed to impede the FSM Department of Justice authority to enforce the nation's

1 criminal or civil laws against any Establishment and/or
2 pharmaceutical activity."

3 Section 25. This act shall become law upon approval by the
4 President of the Federated States of Micronesia or upon its
5 becoming law without such approval.

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9 June 21, 2022

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14 David W. Panuelo
15 President
16 Federated States of Micronesia
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