



REPUBLIC OF KENYA
THE NATIONAL ASSEMBLY
THIRTEENTH PARLIAMENT – SECOND SESSION – 2023
DIRECTORATE OF DEPARTMENTAL COMMITTEES
DEPARTMENTAL COMMITTEE ON HEALTH

REPORT OF
THE DEPARTMENTAL COMMITTEE ON HEALTH
ON
THE RATIFICATION OF AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF
THE AFRICAN MEDICINES AGENCY (AMA)

Directorate of Departmental Committees,
Clerk's Chambers,
Parliament Buildings,
NAIROBI.


 THE NATIONAL ASSEMBLY PAPERS LAID	
DATE: 22 MAR 2023	DAY: Wednesday
TABLED BY:	Hon. Patrick Munene Vice Chair, Health Committee
CLERK-AT THE-TABLE:	Anne Shibuko
	March, 2023

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CHAIRPERSON'S FOREWORD

The Cabinet Secretary, Ministry of Foreign Affairs, submitted a memorandum to the National Assembly dated 18th May, 2022 regarding the ratification of the African Union Treaty for the establishment of the African Medicines Agency (AMA). The memorandum and text of the Protocols were committed to the Departmental Committee on Health for processing. Considering that, the House proceeded to *Sine die* recess immediately thereafter, marking the end of the 12th Parliament, the paper could not be considered. The aid treaty was re-tabled before the House on Thursday, December 1, 2022 in the 13th Parliament.

The African Union treaty on establishment of the African Medicines Agency (AMA) was approved on 12th May, 2022 by Cabinet during its meeting. Considering the protocols, the Committee held a total of five sittings.

Pursuant to the provisions of Article 118 (1)(b) of the Constitution on public participation and section 8(3) of the Treaty Making and Ratification Act of 2012, the Committee placed advertisements in two local dailies of nationwide circulation, on 26th January 2023 requesting for submissions of memoranda on the subject. The Committee received a memorandum in support of the African Medicines Agency (AMA).

Further, the Committee deliberated on the treaty with the agencies involved, in recognition of the crosscutting nature of the treaty.

The Committee is thankful to the Office of the Speaker and the Clerk of the National Assembly for the logistical and technical support accorded to it during its Sittings.

Pursuant to Section 8(4) of the Treaty Making and Ratification Act, 2012 and Standing Order 199, it is my pleasant duty to present the Report of the Departmental Committee on Health on its consideration of the treaty on the establishment of the African Medicines Agency (AMA).

**HON. DR. ROBERT PUKOSE, MP- CHAIRPERSON
DEPARTMENTAL COMMITTEE ON HEALTH**

1.0 PREFACE

1.1 Establishment of the Committee

The Departmental Committee on Health is established pursuant to Standing Order 216.

1.2 Mandate of the Committee

The Committee is mandated under Standing Order 216 (4) and (5) to inter alia-

- a) *investigate, inquire into, and report on all matters relating to the mandate, management, activities, administration, operations and estimates of the assigned ministries and departments;*
- b) *study the programme and policy objectives of ministries and departments and the effectiveness of the implementation and effectiveness of the implementation;*
- c) *study and review all legislation referred to it;*
- d) *study, assess and analyze the relative success of the ministries and departments as measured by the results obtained as compared with their stated objectives;*
- e) *investigate and inquire into all matters relating to the assigned ministries and departments as they may deem necessary, and as may be referred to them by the House;*
- f) *vet and report on all appointments where the Constitution or any law requires the National Assembly to approve, except those under Standing Order 204 (Committee on Appointments);*
- g) ***examine treaties, agreements and conventions;***
- h) *make reports and recommendations to the House as often as possible, including recommendation of proposed legislation;*
- i) *consider reports of Commissions and independent offices submitted to the house pursuant to the provisions of Article 254 of the Constitution; and*
- j) *examine any questions raised by Members on a matter within its mandate.*

In executing its mandate, the Committee oversees the Ministry of Health;

According to second Schedule of the Standing Orders, the Committee is mandated to consider the following subjects:

- i. Health;
- ii. Medical care and Health insurance including universal health coverage.

1.3 Committee Membership

The Committee comprises the following fifteen (15) Members;

4. The Committee was constituted by the House on 27th October 2022 and comprises the following Members;

Chairperson

Hon. (Dr.) Robert Pukose, MP
Endebes Constituency
UDA Party

Vice-Chairperson

Hon. Ntwiga, Patrick Munene MP
Chuka/Igambang'ombe Constituency
UDA Party

Members

Hon. Owino Martin Peters, MP
Ndthiwa Constituency
ODM Party

Hon. Julius Ole Sunkuli Lekakeny, MP
Kilgoris Constituency
KANU

Hon. Muge Cynthia Jepkosgei, MP
Nandi (CWR)
UDA Party

Hon. Maingi Mary, MP
Mwea Constituency
UDA Party

Hon. Wanyonyi Martin Pepela, MP
Webuye East Constituency
Ford Kenya Party

Hon. Mathenge Duncan Maina, MP
Nyeri Town Constituency
UDA Party

Hon. Kipngok Reuben Kiborek, MP
Mogotio Constituency
UDA Party

Hon. Lenguris Pauline, MP
Samburu (CWR)
UDA Party

Hon. Nyikal James Wambura, MP
Seme Constituency
ODM Party

Hon. Oron Joshua Odongo, MP
Kisumu Central Constituency
ODM Party

Hon. Kibagendi Antoney, MP
Kitutu Chache South Constituency
ODM Party

Hon. (Prof.) Jaldesa Guyo Waqo
Moyale Constituency
UPIA Party

Hon. Mukhwana Titus Khamala, MP
Lurambi Constituency
ANC Party

1.4 Committee Secretariat

1. The following are the Secretariat who support the Committee;

Mr. Hassan Abdullahi Arale
Clerk Assistant II/Head of Secretariat

Mr. Gladys Jepkoech Kiprotich
Clerk Assistant III

Ms. Marlene Ayiro
Principal Legal Counsel II

Ms. Salat Abdi Ali
Senior Serjeant-At-Arms

Ms. Faith Chepkemoi
Legal Counsel II

Mr. Yakub Ahmed
Media Relations Officer II

Mr. Rahab Chepkilim
Audio Recording Officer II

Ms. Abigel Muendi
Research Officer III

Mr. Hiram Kimuhu
Fiscal Analyst III

Mr. Benson Kimanzi
Serjeant-At-Arms III

2.0 ANALYSIS OF THE AGREEMENT

INTRODUCTION

1. Article 2(5) of the Constitution of Kenya, 2010 provides that the general rules of international law while Article 2(6) of the Constitution provides that any treaty or convention ratified by Kenya shall form part of the law of Kenya under this Constitution.
2. The Treaty Making and Ratification Act, No. 45 of 2012 (hereinafter referred to as “the Act”) was enacted by Parliament to give effect to Article 2(6) of the Constitution. The Act governs the making and ratification of treaties in Kenya.
3. Section 2 of the Act defines a treaty as an “international agreement concluded between States in written form and governed by international law, whether embodied in a single instrument or in two or more related instruments and whatever its particular designation and includes a convention”.
4. Under the Constitution and the Act, the responsibility of initiating the treaty making process, negotiating and ratifying a Treaty lies with the Executive. In making this decision, the Executive ought to be guided by Section 5(2) of the Act which provides considerations that must be followed including:
 - a) the need that the new treaty is to meet;
 - b) the existing legal regime, including the extent of its applicability to the perceived problem;
 - c) the probability of reaching the required measure of agreement on the solution aimed for;
 - d) any relevant legislative efforts related to the perceived problem;
 - e) the optimal form for the proposed treaty;
 - f) the likelihood that the proposed treaty shall be accepted by a sufficient number of states, where the treaty is multilateral;
 - g) the anticipated time schedule for completing the treaty-making process;
 - h) the expected costs of formulating and adopting the treaty to Kenya; and
 - i) in formulating treaties relating to technical or scientific problems; whether extensive scientific studies or research have been carried out to determine the parameters of the problem and the lines of potential solutions.

ROLE OF THE NATIONAL ASSEMBLY IN TREATY MAKING AND RATIFICATION

5. Although initiation of the treaty making process is the role of the Executive, Parliament as the legislative arm decides whether a Treaty shall form part of the law of Kenya upon which the treaty comes into force. This flows from Article 94(5) of the Constitution which provides that “no person or body, other than Parliament, has the power to make provision having the force of law in Kenya except under authority conferred by this Constitution or by legislation”.
6. After the Treaty has been approved by the National Assembly, it therefore becomes binding upon Kenya and Kenya cannot invoke the provisions of its domestic law to justify any failure to perform its obligations under a treaty ratified by it.

7. According to the Vienna Convention on the Law of Treaties, 1969 which governs the making and ratification of treaties internationally, a treaty becomes binding on a state upon ratification.
8. Section 2 of the Treaty Making and Ratification Act defines ratification as the “the international act by which the State signifies its consent to be bound by a treaty and includes acceptance, approval and accession where the treaty so provides”.
9. Under section 7 of the Act, where the Government intends to ratify a treaty, the Cabinet Secretary of the relevant State department shall, in consultation with the Attorney-General, submit to the Cabinet the treaty, together with a memorandum outlining—
 - a) the objects and subject matter of the treaty;
 - b) any constitutional implications including—
 - i. any proposed amendment to the Constitution; and
 - ii. that the treaty is consistent with the Constitution and promotes constitutional values and objectives;
 - c) the national interests which may be affected by the ratification of the treaty;
 - d) obligations imposed on Kenya by the treaty;
 - e) requirements for implementation of the treaty;
 - f) policy and legislative considerations;
 - g) financial implications;
 - h) ministerial responsibility;
 - i) implications on matters relating to counties;
 - j) the summary of the process leading to the adoption of the treaty;
 - k) the date of signature;
 - l) the number of states that are party to the treaty;
 - m) the views of the public on the ratification of the treaty;
 - n) whether the treaty sought to be ratified permits reservations and any recommendations on reservations and declarations;
 - o) the proposed text of any reservations that should be entered when ratifying the treaty in order to protect or advance national interests or ensure conformity with the Constitution; and
 - p) whether expenditure of public funds will be incurred in implementing the treaty and an estimate, where possible, of the expenditure.

Consideration by the National Assembly

The Treaty Making and Ratification Act, No. 45 of 2012

10. Section 8 of the Treaty Making and Ratification Act, No. 45 of 2012 provides for the consideration of Treaties by Parliament. Upon approval of a Treaty by Cabinet, the relevant Cabinet Secretary shall submit the Treaty together with a memorandum on the Treaty to the Speaker of the National Assembly for tabling pursuant to the Standing Orders.
11. Section 8(3) of the Treaty Making and Ratification Act, No. 45 of 2012 provides that the relevant parliamentary Committee in the National Assembly is tasked with consideration of the Treaty and shall ensure public participation in the ratification process in accordance with the laid down parliamentary procedures. (Section 8(3) of the Act).

Decision on Ratification by the National Assembly

12. The National Assembly may:
- a) **refuse to approve the ratification of a Treaty**-where the National Assembly refuses to approve the ratification of a treaty, the Clerk of the National Assembly shall submit the resolution of the House to the relevant Cabinet Secretary within fourteen (14) days of such resolution (Section 8(7) of the Act) and the Government shall not ratify the said Treaty;
 - b) **approve the ratification of a Treaty without reservations to specific provisions of the treaty** (Section 8(4) of the Act)-where the ratification of a treaty is approved by National Assembly without any reservations to the treaty, the relevant Cabinet Secretary (the Cabinet Secretary for the time being responsible the subject matter of the treaty) shall, within thirty (30) days from the date of the approval of the ratification of treaty request the Cabinet Secretary to prepare the instrument of ratification of the treaty;
 - c) **approve the ratification of a Treaty with reservations to specific provisions of the treaty**-where a treaty is approved for ratification with reservations to some provisions of the treaty, the treaty shall be ratified with those reservations to the corresponding article in the treaty.
13. Proposed reservations made by the National Assembly are introduced as a provision into the Treaty in line with the procedure set out in the Standing Orders (Section 8(5) of the Act).
14. In making the decision on the approval for ratification of a Treaty, Section 8(9) of the Act provides that the National Assembly shall not approve:
- a) the ratification of a treaty or part of it if its provisions are contrary to Constitution; and
 - b) a reservation to a treaty or part of it if that reservation negates any of the provisions of the Constitution even if the reservation is permitted under the relevant treaty.
15. Section 12 of the Act provides that a Treaty cannot be ratified unless the same has been considered and approved by the Cabinet and Parliament. A person who ratifies a Treaty without following this process commits an offence and shall be liable to imprisonment for a term not exceeding fifteen (15) years or to a fine not exceeding twenty (20) million shillings or to both such fine or imprisonment.

1. The National Assembly Standing Orders

16. One of the functions of Departmental Committees under Standing Order 216(5)(fa) is to *“examine treaties, agreements and conventions”*.
17. The procedure of ratification of treaties is guided by Part XXI and in particular Standing Order 170A of the National Assembly Standing Orders. Standing Order 170A provides:
- “(1) A treaty submitted to the National Assembly for ratification shall be laid on the Table of the House and stand committed to the relevant Committee for consideration.*
- (2) The committee shall undertake public participation before submitting its report to the House.*

(3) In addition to the information required to be submitted to the National Assembly under written law, the committee may require the relevant Cabinet Secretary to submit further information, including—

a) the social and environmental impact of the treaty in the short-term, medium term and long-term; and,

b) the nature and evidence of any public participation conducted on the treaty.

(4) The report of the committee to the House shall include—

a) information on the views of the people on the ratification of the treaty emanating from public participation conducted by the committee;

b) the findings of the committee on the treaty and any other information the committee may deem necessary; and

c) a recommendation that the House—

(i) approves the ratification of the treaty, or

(ii) approves the ratification of the treaty with reservations, or

(iii) rejects the ratification of the treaty.

(5) In approving ratification of a Treaty with reservations, the House shall specify the affected provisions of the Treaty and the proposed text of each reservation, which may include prescription of timelines within which an obligation is to be fulfilled before implementation of the Treaty.

(6) Upon decision of the House on a Treaty, the Clerk shall, within seven (7) days, notify the relevant Cabinet Secretary and enter the information in the register of treaties.”

A. OBJECTIVE OF THE TREATY

18. The African Union (AU) Treaty for the Establishment of the African Medicines Agency (hereinafter “the Treaty”) was adopted by the 32nd ordinary session decision of the Assembly of Heads of State and Government on 11th February 2019.

19. The Treaty establishes the African Medicines Agency (AMA) under Article 3. AMA is a specialized agency of the AU with its own rules, membership and resources, intended to enhance the capacity of state parties and Regional Economic Communities (RECs) to regulate medical products in order to improve access to quality, safe and efficacious medical products on the African continent.

20. Weak regulatory systems have resulted in the circulation of substandard and falsified medical products in many African Union member states causing risk to public health, harm to patients and undermining confidence in healthcare delivery systems. The AMA therefore intends to:

a) Provide a platform for coordination and strengthening of on-going regional and continental harmonization initiatives

b) Complement efforts of RECs and contribute to their capacity building towards improving access to quality assured medical products with the agenda of Universal Health Coverage and Sustainable Development Goals

- c) Define acceptable standards in the regulation of medical products in the continent

21. The Organs of the AMA are:

- (a) The Conference of the State Parties-the highest policy-making organ of the Agency. It is composed of all member states of the African Union (AU) who ratify the Treaty and which will be represented by their Ministers responsible for health or their representatives. The conference shall meet once every two (2) years.
- (b) Governing Board-it shall be composed of the heads of National Medicines Regulatory Authorities (NMRAs), RECs, Regional Health Organizations responsible for regulatory affairs among others.
- (c) The Secretariat-responsible for coordinating the implementation of the decisions of the Conference of State Parties and Policy organs of the AU and the Board of the AMA. The secretariat shall be headed by the Director General who shall be responsible for the day-to-day management of the AMA.
- (d) The Technical Committees-The Board shall permanent and ad hoc technical committee to provide technical guidance on specific areas of regulatory expertise.

22. The obligations of State Parties under the Treaty include:

- a) To coordinate national and sub-regional medicines regulatory systems;
- b) To conduct regulatory oversight of selected medical products including traditional medicines;
- c) To promote cooperation, harmonization and mutual recognition of regulatory decision;
- d) To strength and harmonize efforts of the AU-recognized RECs, Regional Health Organizations (RHOs) and Member states; and
- e) To complement and enhance collaboration and contribute to improving patient's access to quality, safe and efficacious medical products and health technologies on the continent.

23. The AMA is supposed to work closely with the AU, World Health Organization (WHO), African Centres for Disease Control and Prevention (Africa CDC), and any other UN agencies. It shall further maintain active cooperation with AU member states and other countries as well.

24. Article 33 allows a State Party when ratifying the Treaty to submit reservations to any provisions of the Treaty in writing. The reservation should not contravene the objects and purpose of the Treaty. The reservation may be withdrawn at any time in writing.

25. Article 34 allows a State Party to withdraw from the Treaty three (3) years from the date of entry into force of the Treaty provided that the obligations of such a party prior to the withdrawal shall still subsist.

26. The Treaty may be dissolved by an agreement of two-thirds of the State Parties to the Treaty and may be amended or revised pursuant to Article 35 and 36 of the Treaty.

27. Under Article 37, the Treaty is open for signature and ratification by Member Sates of the AU.

28. Under Article 39 of the Treaty, the Treaty shall enter into force thirty (30) days after deposit of the fifteenth (15th) instrument of ratification. For countries such as Kenya that are ratifying the Treaty after it has come into force, the Treaty shall come into force on the date of deposit of instrument of accession or ratification.

B. PUBLIC PARTICIPATION ON THE TREATY

a) Legal Provision on Public Participation

29. Article 118 (1) (b) of the Constitution of Kenya provides as follows
“Parliament shall facilitate public participation and involvement in the legislative and other business of Parliament and its Committees.”
30. Section 8 of the Treaty Making and Ratification Act, No. 45 of 2012 provides for the consideration of Treaties by Parliament. Upon approval of a Treaty by Cabinet, the relevant Cabinet Secretary shall submit the Treaty together with a memorandum on the Treaty to the Speaker of the National Assembly for tabling pursuant to the Standing Orders.
31. Section 8(3) of the Treaty Making and Ratification Act, No. 45 of 2012 provides that:
*“the relevant parliamentary Committee shall, during its consideration of the Treaty, ensure **public participation** in the ratification process in accordance with laid down parliamentary procedures”.*
32. Standing Order 170A provides:
*“(2) **The committee shall undertake public participation before submitting its report to the House.***
(4) The report of the committee to the House shall include—
d) information on the views of the people on the ratification of the treaty emanating from public participation conducted by the committee;

(b) Methodology used by the Committee in Public Participation

33. The Memorandum by the Ministry of Foreign Affairs on the Ratification of the African Union Treaty for the Establishment of the African Medicines Agency (AMA) was laid on the Table of the House on Tuesday, 7th June 2022. The Treaty was however not considered as the House in the 12th Parliament proceeded to *Sine die recess* immediately thereafter.
34. The Treaty was re-tabled before the House on Thursday, 1st December 2022 in the 13th Parliament and committed to the Departmental Committee on Health for consideration.
35. Pursuant to the aforementioned provisions of the Constitution, the Treaty Making and Ratification Act, 2012 and Standing Orders, the Committee through local daily newspapers of 26th January, 2023 published an advertisement inviting the public to submit memoranda. Further, in a letter dated 25th January, 2023, the Committee wrote to various stakeholders including the Ministry of Foreign Affairs, National Treasury, Ministry of Health, Ministry of Trade, Investment and Industry, Ministry of East African

Community, Office of the Attorney General and Department of Justice, Kenya Revenue Authority, Kenya Law Reform Commission to submit memorandum on the Treaty which they all supported the treaty (*responses attached*).

36. The Committee also held a stakeholder engagement forum on 27th February 2023 with various non-state actors and non-governmental organizations at Mercure Hotel, Nairobi.

The stakeholders who attended the forum were:

- (a) ROCHE
- (b) Coalition for Health Research and Development (CHREAD)
- (c) PATH
- (d) Kenya Pharmaceutical Association
- (e) International AIDS Vaccine Initiative (IAVI)
- (f) DNDI
- (g) Generic Specialities
- (h) Federation of Kenya Pharmaceutical Manufacturers
- (i) Pharmaceutical Society of Kenya
- (j) Renal Patients Society of Kenya
- (k) NCD Alliance of Kenya
- (l) Kenya Medical Laboratory Technicians and Technologist Board
- (m) Mission for Essential Drugs and Supplies (MEDS)
- (n) Ministry of Health, Directorate of Health Product and Technologies
- (o) United States Pharmacopeia (USP)
- (p) Pharmacy and Poisons Board
- (q) National Quality Control Laboratory
- (r) African Medical and Research Foundation (AMREF)
- (s) MI-PH

37. The report is divided into two parts as follows:

Part I of the Report contains the analysis of the public submissions on the ratification of the Treaty, written and oral submissions received from the public and various stakeholders noting general comments in support or against the ratification of the Treaty and the list of institutions that submitted their memoranda.

38. Part II of the Report contains a copy of the newspaper advertisements of Wednesday, 26th January, 2023 inviting the public to submit memoranda on the ratification of the Treaty and a letter inviting the relevant stakeholders for memoranda and the minutes of the Committee sittings during the consideration of the ratification of the Treaty.

3.0 STAKEHOLDER VIEWS ON THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY

39. The table below highlights the stakeholder comments on the ratification of the Treaty—

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)			
	STAKEHOLDER	POSITION	JUSTIFICATION
1.	State Department for East African Community Ministry of East African Community and Regional Development	After consultations with the Ministry of Health, supports ratification of the Treaty vide letter dated 2 nd February 2023.	<ul style="list-style-type: none"> • AMA flows from the African Medicines Regulatory Harmonization (AMRH) Initiative that has been advanced by the AU Development Agency and regional economic communities (RECs) in collaboration with development partners. • Kenya supported AMRH initiatives- set up of EAC-MRH Programme and implementation-helped Kenya realize its health sector development goals • AMA to complement efforts of existing national and regional regulatory bodies or harmonization initiatives at RECs level which will continue with their work • Kenya to benefit when the treaty comes into force as follows: <ol style="list-style-type: none"> a) AMA provides platform for coordination and strengthening on-going regional and continental harmonization initiatives, pool expertise and capacities for optimal use of the limited resources and combat of substandard and falsified medical products b) Will strengthen Kenya's clinical trials ecosystem including Covid-19, manufacturing industry, and ability to regulate and monitor safety of health products
2.	Ministry of Foreign and Diaspora Affairs Office of the Attorney General Ministry of Health	Jointly supports the ratification of the Treaty vide letter dated 3 rd February 2023. No reservations raised.	<ul style="list-style-type: none"> • MOH through the Pharmacy and Poisons Board (PPB) ensures quality, safety and efficacy of health products and technologies through capacity building, WHO collaborative procedures, harmonization initiatives, collaboration with development partners • Kenya through PPB has been contributing technically to the AMRH initiatives that has facilitated the realization of Kenya's health

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<p>sector development goals; Two (2) officers of PPB are Chairpersons of Technical Committees under the AUDA-NEPAD AMRH Initiative, the precursor of the AMA</p> <ul style="list-style-type: none"> • AMA headquarters in EAC (Rwanda) • 23-member states have ratified Treaty (Rwanda and Uganda have fully ratified and deposited instruments to the AU Commission) • Covid-19 triggered the interest of African countries to develop their manufacturing capacities to remedy challenge of access to essential health products such as vaccines when global supply chains deprioritize Africa's needs • Depicts Kenya's commitment to Africa's collective action for improved regulation of medicines, medical products and technologies as a Member of AU and RECs such as EAC and IGAD • Ratification facilitates the achievement of Kenya Vision 2030 and supports the achievement of objectives under the Kenya National Health Policy, Kenya Health Sector Strategic Plan and Kenya National Pharmaceutical Policy. • Kenya has the largest pharmaceutical industry in the common market for the Eastern and Southern African Regions- over 30 manufacturing plants • AMA will open up market for Kenyan pharmaceutical products from USD 160 Million (EAC) to USD 1.2 Billion (African Continent) <p><i>Legal Implication</i></p> <ul style="list-style-type: none"> • Treaty in line with/ not amending the Constitution • May require amendment of Kenyan laws for compliance with and implementation of the Treaty obligations-Treaty advocates for adoption of the AU model law on regulation of medical products • Kenya may need to develop guidelines for periodic reporting obligations

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			arising from joint capacity assessments for Member States
3.	Pharmaceutical Society of Kenya (PSK)-Professional Body of Pharmacists in Kenya)	Support the ratification of the Treaty vide letter dated 8 th February 2023.	<ul style="list-style-type: none"> • Given that the objectives of the Treaty on Medicines entail ensuring public safety, promoting innovation, streamlining product registration, harmonizing healthcare education systems among others, Pharmacists are important stakeholders in this and their role and contribution should be given more prominence in the entire process of establishing the AMA. Their expertise can be applied to several areas such as innovation of new therapeutic products and systems technologies and management of non-communicable diseases. They can form a think tank funded by AU or AMA to collect data and offer policy direction and guidance on effective use of medicines. They can also work with higher learning regulators in developing a harmonized curriculum for pharmacists in the continent. • The Treaty encourages local manufacturing. • In relation to resourcing, the Treaty is presently being funded by donors, what will happen later when donors pull out? • The role of pharmacists and the pharmaceutical profession needs to be leveraged upon in the implementation of the Treaty.
4.	Kenya AIDS NGOs Consortium (KANCO)	Supports the expedited ratification of the Treaty vide letter dated 9 th February 2023.	<ul style="list-style-type: none"> • The Treaty provides: <ul style="list-style-type: none"> ✓ Support for growth of local pharmaceutical production ✓ Mechanism for evaluating medical products for treatment of priority diseases ✓ Coordination of joint reviews of clinical trial applications for vaccines ✓ Information sharing and collaboration with RECs and National Medicines Regulatory Authorities in identification of

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<p>substandard and falsified medical products</p> <ul style="list-style-type: none"> • Kenya is a major player in the field of global health should be in the forefront in ratifying the Treaty which will promote Article 43 of Constitution on the right to access the highest attainable standard of health including reproductive health care. • The Treaty has been ratified by 33 out of 55 AU Member States. • 20 nations have ratified and deposited the Treaty; 3 have ratified but not deposited, 22 not ratified or signed, 10 including Kenya have signed but not ratified. • There is need to establish an African Digital Platform that lists the approved medical products and technologies in the Continent. • The Association applauded the Kenyan Parliament for undertaking public participation on Treaty.
5.	<p>Coalition for Health Research and Development</p> <p>Note: KANCO is part of this coalition.</p>	<p>Supports the expedited ratification of the Treaty vide letter dated 9th February 2023</p>	<ul style="list-style-type: none"> • Ratification of the Treaty will catalyze realization of Universal Health Coverage through faster access to the highest quality of medical products thereby facilitating the realization of Article 43 of the Constitution. • The Coalition supports harmonization of regulatory systems for medical products in the country. • The Treaty is in line with the Constitution and will contribute to achievement of government health policies such as Kenya Health Policy, Kenya National Pharmaceutical Policy etc. • Ratification will lead to job creation, economic growth, reduced over reliance in imported expensive medicines and medical products and increased local manufacturing capabilities.

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<ul style="list-style-type: none"> • AMA will provide capacity building upon request, to national regulatory authorities such as the Pharmacy and Poisons Board. • After ratification, there is need to identify gaps and strengthen the existing legislation. • There is need to assess elements of patient safety so as to reduce harm to consumers and ensure the intended therapeutic outcomes of the Treaty.
6.	Kenya Revenue Authority	Supports the ratification of the Treaty vide letter dated 13 th February 2023.	KRA does not have additional input on the Treaty and Memorandum from the Ministry of Foreign Affairs.
7.	Office of the Attorney General and Department of Justice	Supports the ratification of the Treaty vide letter dated 2 nd February 2023.	The Treaty and Memorandum from the Ministry of Foreign Affairs are in order from a legal perspective.
8.	Federation of Kenya Pharmaceutical Manufacturers	Supports the ratification of the Treaty vide letter dated 1 st March 2023.	<ul style="list-style-type: none"> • How will all African pharmaceutical companies have a level playing field in protected countries such as Ghana, and Algeria and Morocco which restricts the list of products that can be imported into the country and support sale of locally manufactured products? • What mechanisms have been put in place to ensure that AMA will work with individual state parties and RECs in regulation of medical products in the continent? • Kenya should be represented in the AMA Secretariat. • What is the situation with regard to the EAC Harmonization policy? Will money be earned from registration and authorization of medicines? • Involvement of pharmacists in research on medicines- There is need to have a harmonized way of training pharmacists

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<ul style="list-style-type: none"> • There is need to ensure proper disposal of medication • There is fear that the pharmaceutical market might be swamped • There is need to reduce the cost of production so as to improve productivity and cost effectiveness • The Treaty is aligned to the Constitution and policy framework. It makes economic sense and strengthens regulatory expertise and ensures harmonized processes • There is need to ensure Kenya can manufacture and benefit from the sale of Active Pharmaceutical Ingredients (APIs)?
9.	PATH.	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • Treaty ratified on 11th January 2019 and came into force on 5th November 2022. • 23 member countries have fully ratified the Treaty; 10 member countries have signed the Treaty but not completed the ratification process. Kenya part of the 10 countries. • The main objective of Treaty is to enhance capacity of state parties and Regional Economic Communities (RECs). • The Treaty facilitates the implementation of the AU Model Law on Regulation of Medical Products. • AMA will be an advisory institution and a platform for collaboration between national medicines regulatory authorities/ RECs steered by state parties. • The AMA Governing Council to be formed in two weeks' time. Kenya lost the opportunity to host AMA and therefore needs to position its people for the positions of influence in the AMA Governing Council. • The AU Model Law on the Regulation of Medicines is being implemented through the Kenya Drugs Authority Bill

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			sponsored by the Chairperson of the Committee. The Bill also anchors the AMA. The Bill ought to be enacted to guarantee the safety of drugs in Kenya.
10	African Medical and Research Foundation (AMREF)	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • The Treaty ensures access to medicine. • Kenya is the lead in Africa in the pharmaceutical manufacturing sector as it currently manufactures both for the local and external markets. 70% of Kenyan pharmaceutical products are exported • The Treaty encourages capacity building through knowledge transfer and training.
11	Mission for Essential Drugs and Supplies (MEDS)	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • MEDS supports ratification of the Treaty as it will reduce bureaucracy in manufacturing of health products and technologies due to improvement in regulatory requirements • Kenya needs to leverage on the local capacity to ensure quality control • There is need to do post medical surveillance • There is need to fast-track the Kenya Development Authority Bill • There is need to revive the Traditional Practice Bill that was done during the previous Parliament.
12	Renal Patients Society of Kenya and Non-Communicable Diseases (NCD) Alliance of Kenya	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • The Treaty will help in reduction in the cost of drugs and ensure availability of drugs. There is need to ensure local or regional production of drugs using locally available materials and expertise so that drugs can be sold at a cheapest price. This will solve the problem of expensive drugs while ensuring safety and quality. The AMA to provide for a means of capacity building within the continent • Africa was considered last in the supply of vaccines during the covid-19 period.

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
13	Drugs for Neglected Diseases	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • Accelerated access to treatment should focus on policy adoption- will there be reliance on WHO for recommendation before adoption of medicines in the continent? • In relation to the regulatory aspect, does establishment of AMA eliminate the WHO? • What is the effect on customs within the respective jurisdictions? Each and every jurisdiction has to inspect a drug manufacturer's factory to check quality and standards?
14	ROCHE	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • The Treaty will ensure regulatory harmonization and convergence and tackle the following challenges <ul style="list-style-type: none"> (a) Delay in product registration which affects market authorization certification as well as product retention in the market- The EAC takes 90 to 180 days to authorize the sale of drugs within the EAC Member States. (b) Lengthy WHO pre-qualification processes which lengthens the provision of approvals-it takes at least four (4) years to get such pre-qualification. (c) Trade barriers- The Treaty incorporates best practices by providing mechanisms for inspection and audit of drugs. It also links trade and regulation by removing the multiplicity of regulation for instance Kenya has Kenyan Bureau of Standards (KEBS) and Anti-Counterfeit Authority.
15	Kenya Pharmaceutical Association	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • The Association supports the treaty and memorandum by the Ministry of Foreign Affairs. • The focus of Treaty is on governance structure. There is little on practice and the access to health care. How will HRH/healthcare workers benefit from the Treaty? Is there provision for Cross-border practice? Will there be standardization of HRH? What are the norms for HRH that will ensure

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<p>harmonization of training and certification of health care professionals within the continent?</p> <ul style="list-style-type: none"> • Some countries have zero regulations. What will be the effect of the lack of domestic regulations particularly with border entry points restrictions? How does the Treaty guarantee the authenticity of health products? • How are middle level practitioners taken care of under the Treaty?
16	Pharmacy and Poisons Board	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> • The Board supports the Treaty and was part of its development process. • The Treaty will deal with the high cost of manufacturing of medicines and particularly APIs. South Africa and DRC have been extracting quinine. China is the biggest producer of APIs in the country. • There is need to do capacity building within the country for buy in within and outside the country. • There is a huge market for the country as AMA is continental. Manufacturers can distribute drugs within the whole continent and manufacturers can come and set up their businesses in the country. There is therefore need to look at the cost of electricity and taxation so as to encourage the influx of pharmaceutical manufacturers. • There is need to critically assess the therapeutic outcome of the Treaty and tap into it as a country. • There is multiplicity of Regulations on drugs within the country. There is need to avoid duplicity for instance the country has inspectors from PPB and the Anti-Counterfeit Authority have the same responsibilities? The lack of harmonization of laws is the main reason why Kenya is at the maturity level 1. This issue to be addressed in the Kenya

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			<p>Drugs Authority Bill sponsored by the Chairperson of the Committee.</p> <ul style="list-style-type: none"> • The Treaty ensures short marketing timelines and will lead to revenue generation as the country can sell drugs across the continent. • There is a big challenge of capacity in the regulatory entities. The number of people in regulatory authorities is currently less than 15 in number when optimal ought to be 100. • The implementation of the Treaty requires digitization of processes and systems. • The WHO prequalification process only assesses a class of drugs namely malaria, ARVs, Reproductive health, diarrhea in children, Covid-19 and other emergency issues. The Treaty will not replace the WHO but will work alongside WHO for strengthening of capacity within the country and the region. There are assessors working for WHO in Tanzania, Ghana and Kenya. Kenya has two WHO qualified assessors. • There is need to check the ingress of substandard and falsified medicines while checking market control in line with the Head of Public Service circular on border management. • Cross border trading will benefit from the Treaty as there is harmonized regulation. There is however need to build an alert system. • The EAC is working on regulation of pharmacists which will ensure access to quality products • The list of OTC poisons in part I and Part II of the Schedule to the Pharmacy and Poisons Act, Cap. 244. The list needs to be revised. The law also needs to be amended to provide for scheduling and rescheduling of medicines. • There are three categories of drugs namely OTC, prescription only and

THE AFRICAN UNION (AU) TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

	STAKEHOLDER	POSITION	JUSTIFICATION
			pharmacy only. The prescription category has more categories and the PPB has developed some guidelines on the same.
17	Kenya Medical Laboratory Technicians and Technologist Board	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> The Treaty will ensure access to safe and quality health products and technologies in the continent.
18	National Quality Control Lab	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> There is need to develop an alert system for substandard and falsified medicines in the continent. The Treaty should provide for the regulation of herbal products as the Treaty currently only provides for dietary supplements and traditional medicine.
19	Cabinet Secretary, Ministry of Health	Supports the ratification of the Treaty.	<ul style="list-style-type: none"> The African continent was the last to receive the covid-19 vaccines. These vaccines arrived late and many of them expired because of low intake by members of the Public. The Cabinet Secretary requested the Committee to support the Treaty/ World Health Organization (WHO) Logistics Centre in Kenyatta University and the Africa Centre for Disease Control (CDC).

4.0 COMMITTEE OBSERVATIONS

39. The Committee having considered the ratification of the African Union Treaty for the Establishment of the African Medicines Agency and submissions from stakeholders makes the following observations:

- (i) The Cabinet Secretary, Ministry of Foreign and Diaspora Affairs, Dr. Alfred Mutua signed the Treaty on 16th February, 2023;
- (ii) The Treaty is in line with the Constitution of Kenya, 2010 and complies with the provisions of the Treaty Making and Ratification Act, No. 45 of 2012;
- (iii) The Treaty may require amendment of Kenyan laws for compliance with and implementation of the Treaty obligations as the Treaty advocates for the adoption of the African Union Model Law on Regulation of Medical products;
- (iv) Ratification of the Treaty will catalyze realization of Universal Health Coverage through faster access to the highest quality of medical products thereby facilitating the realization of Article 43 of the Constitution;
- (v) The Treaty will contribute to achievement of government health policies such as Kenya Health Policy, Kenya National Pharmaceutical Policy among others; and
- (vi) The Treaty will benefit Kenya as it supports local pharmaceutical production and strengthens the country's ability to regulate and monitor safety of health products.

5.0 FINDINGS

40. Pursuant to the analysis of the submissions and documents tabled, the Committee finds that the African Union Treaty on the establishment of the African Medicines Agency is consistent with constitution and do not propose any amendment to the Constitution.


6.0 COMMITTEE RECOMMENDATION

41. The Committee recommends: -


THAT, Pursuant to Section 8 of the Treaty Making and Ratification Act, the House **APPROVES the Ratification** of the African Union Treaty for the Establishment of the **African Medicines Agency**.

Justification

The African Union Treaty for the Establishment of the African Medicines Agency facilitates the realization of Universal Health Coverage through improved regulation of medical products.

for Signed.....  Date: *16/3/2023*.....

**HON. DR. ROBERT PUKOSE, MP – CHAIRPERSON
DEPARTMENTAL COMMITTEE ON HEALTH**

 THE NATIONAL ASSEMBLY PAPERS LAID	
DATE: <i>22 MAR 2023</i>	DAY: <i>Wednesday</i>
TABLED BY:	<i>Hon. Patrick Munene Vice Chair, Health Committee</i>
CLERK-AT THE-TABLE:	

ADOPTION LIST OF THE MEMBERS



THE NATIONAL ASSEMBLY
13TH PARLIAMENT - SECOND SESSION - 2023
DIRECTORATE OF DEPARTMENTAL COMMITTEES
DEPARTMENTAL COMMITTEE ON HEALTH,
ATTENDANCE REGISTER

AGENDA: ADOPTION OF THE REPORT

DATE: 16-02-2023 TIME: From To

VENUE: COMMITTEE ROOM CONTINENTAL 2ND FLOOR

NO.	NAME	SIGNATURE
1.	The Hon. Dr. Pukose Robert, M.P.- Chairperson	
2.	The Hon. Ntwiga Patrick Munene, M.P.-Vice-Chairperson	
3.	The Hon. Maingi Mary, M.P.	
4.	The Hon. Muge Cynthia Jepkosgei, M.P	
5.	The Hon. Kipngor Reuben Kiborek, M.P.	
6.	The Hon. Wanyonyi Martin Pepela, M.P	
7.	The Hon. Mathenge Duncan Maina, M.P.	
8.	The Hon. Lenguris Pauline, M.P.	
9.	The Hon. Oron Joshua Odongo, M.P.	
10.	The Hon. Dr. James Nyikal Wambura, M.P.	
11.	The Hon. Kibagendi Antoney, M.P.	
12.	The Hon. Sunkuli Julius Lekakeny Ole, EGH, EBS M.P.	
13.	The Hon. Prof. Jaldesa Guyo Waqo, M.P.	
14.	The Hon. Titus Khamala, M.P	
15.	The Hon. Owino Martin Peters, M.P.	

Committee Clerk: Hessan A. Arale Signature:

Director, Departmental Committees: Signature:

MINUTES OF THE SITTINGS

MINUTES OF THE THIRTY FIRST (31ST) SITTING OF THE DEPARTMENTAL COMMITTEE ON HEALTH HELD IN COMMITTEE ROOM IN 2ND FLOOR CONTINENTAL HOUSE, PARLIAMENT BUILDINGS ON THURSDAY 16TH MARCH, 2023 AT 10:00 AM.

PRESENT

1. The Hon. Ntwiga Patrick Munene, M.P -**Vice-Chairperson**
2. The Hon. Sunkuli Julius Lekakeny Ole, EGH, EBS, M.P
3. The Hon. Prof. Jaldesa Guyo Waqo, M.P.
4. The Hon. Mary Maingi, MP.
5. The Hon. Lenguris Pauline, M.P
6. The Hon. Oron Joshua Odongo, M.P.
7. The Hon. Mathenge Duncan Maina, M.P
8. The Hon. Owino Martin Peters, M.P.
9. The Hon. Kipngor Reuben Kiborek, M.P

ABSENT WITH APOLOGY

1. The Hon. Dr. Pukose Robert, M.P - **Chairperson.**
2. The Hon. Muge Cynthia Jepkosgei, M.P
3. The Hon. Kibagendi Antony, M.P.
4. The Hon. Titus Khamala, M.P.
5. The Hon. Wanyonyi Martin Pepela, M.P
6. The Hon. Dr. Nyikal James Wambura, M.P.

COMMITTEE SECRETARIAT

1. Mr. Hassan A. Arale - Clerk Assistant II
2. Ms. Abigel Muinde - Research Officer III
3. Mr. Benzon Kimanzi -Serjeant At Arms

MIN. NO. NA/DC-H/2023/132: PRELIMINARIES/INTRODUCTION


The meeting was called to order at 10.30 am with a word of prayer by the Hon. Patrick Ntwiga Munene, M.P. Vice-Chairperson and welcomed members to the meeting.


MIN. NO. NA/DC-H/2023/133: CONSIDERATION AND ADOPTION OF THE REPORT ON THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

The Committee considered the report on the ratification of the African Union Treaty for the establishment of the African Medicines Agency and confirmed and adopted as the true reflection of the Committee deliberations after it was proposed by the Hon. Sunkuli Julius Lekakeny Ole, EGH, EBS, M.P and seconded by the Hon. Prof. Jaldesa Guyo Waqo, M.P.

MIN. NO. NA/DC-H/2023/134: ADJOURNMENT

There being no any other business, The Chairperson, adjourned the meeting at exactly 1.00 p.m.

Sign.......... Date 21/11/23.....

 **HON. DR. ROBERT PUKOSE, M.P – CHAIRPERSON DEPARTMENTAL COMMITTEE ON HEALTH**

MINUTES OF THE FIFTEENTH (15TH) SITTING OF THE DEPARTMENTAL COMMITTEE ON HEALTH HELD IN MERCURE HOTEL IN NAIROBI ON MONDAY 27TH FEBRUARY, 2023 AT 9.30 A.M.

PRESENT

1. The Hon. Dr. Pukose Robert, M.P - **Chairperson.**
2. The Hon. Ntwiga Patrick Munene, M.P -**Vice-Chairperson.**
3. The Hon. Titus Khamala, M.P
4. The Hon. Prof. Jaldesa Guyo Waqo, M.P.
5. The Hon. Wanyonyi Martin Pepela, M.P
6. The Hon. Lenguris Pauline, M.P
7. The Hon. Mary Maingi, MP.
8. The Hon. Dr. Nyikal James Wambura, M.P.
9. The Hon. Oron Joshua Odongo, M.P.
10. The Hon. Muge Cynthia Jepkosgei, M.p

MEMBERS ABSENT WITH APOLOGY

1. The Hon. Kipngor Reuben Kiborek, M.P.
2. The Hon. Mathenge Duncan Maina, M.P
3. The Hon. Kibagendi Antony, M.P.
4. The Hon. Sunkuli Julius Lekakeny Ole, EGH, EBS, M.P
5. The Hon. Owino Martin Peters, M.P.

COMMITTEE SECRETARIAT

1. Mr. Hassan A. Arale - Clerk Assistant II
2. Ms. Gladys Kiprotich - Clerk Assistant III
3. Ms. Faith Chepkemoi -Legal counsel II
4. Ms. Marlene Ayiro -Principal Legal Counsel II
5. Mr. Benson Kimanzi - Serjeant at Arms

STAKEHOLDERS IN ATTENDANCE

1. Mr. Christopher Odera -Regulatory Policy Manager
2. Mr. Simion Bolo -Head of Leis access
3. Mr. Philip Nyakwama - **CHRead**
4. Ms. Mary Muiya - **CHRead**
5. Mr. Peter waiganji - **CHRead**
6. Mr. Johh Paul -**PATH**-Senior Policy Advocacy
7. Mr. Eric Gichare -Kenya Pharmaceutical Associations Secretary General
8. Mr. Ethel Makila - Director for commons advocacy general
9. Ms. Rosemarie Muganda -**RAD, PATH**
10. DR. Julius Machira -General Specialties
11. Dr. Vimal -**FKPM**
12. Mr. John Gikonyo -Renal patients Society of Kenya
13. Mr. Titus Mutwiri -Chairman, Kenya Medical Laboratory Technicians and Technologist
14. Dr. Nur Said - Pharmacist
15. Mr. Shadrack Meme - Health System Specialist
16. Mr. Meshack Ndolo - Manager PH East Africa
17. Mr. Boar Dulloh - Kenya Pharmaceutica I
18. Dr. Pauline Duya -Ministry of Health
19. Dr. Sarah Muteru -**NOCL**

- | | |
|--------------------------|-------------------------------|
| 20. Dr. Sarah Mwangi | -Ministry of Health |
| 21. Mr. Bintiomari Tsala | - KMLTTB- MOH |
| 22. Dr. Juliet Konye | -United States Phacelia (USP) |

MIN. NO. NA/DC-H/2023/65: PRELIMINARIES/INTRODUCTION

The meeting was called to order at 9.30 a.m. with a word of prayer by the Hon. Dr. Pukose Robert, MP Chairperson. Introductions was then done by the members and secretariat followed by different stakeholders.

MIN. NO. NA/DC-H/2023/66: BRIEF BY THE CHAIRPERSON AND THE SENIOR POLICY ADVOCACY PATH ON THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY

a) Brief by chairperson of the committee on health

The Chairman indicated that, he has reintroduced the Kenya Drug Authority Bill which lapsed at the 2nd reading due to sine die recess, and He has removed aspects of blood, cosmetics, food, dieticians, and he is also looking at including the regulation of pharmaceutical practice- training, certification, regulation of training.

b) Brief By the Senior Policy Advocacy Path

He did introduction about AMA and stated that the Treaty was ratified on 11th January 2019; and came into force on 5th November 2022- 23and countries have fully ratified the treaty.

He also stated that the treaty establishes the African Medicines Agency (AMA) as a specialized agency with the objective of enhancing the capacity of State Parties and Regional Economic Communities (RECs) to regulate medical products in order to improve access to quality, safe and efficacious medical products on the African continent. He also added that due to the Weak regulatory systems resulted in the circulation of substandard and falsified (SF) medical products in many African Union member states causing risk to public health, harm to patients and undermining confidence in healthcare delivery systems and AMA was therefore intended to do the following;

1. Provide a platform for coordination and strengthening of on-going regional and continental harmonization initiatives
2. Complement efforts of RECs and contribute to their capacity building towards improving access to quality assured medical products with the agenda of Universal Health Coverage and Sustainable Development Goals.
3. Define acceptable standards in the regulation of medical products in the continent

He further highlighted that, some Donor funding-at the beginning of implementation of the treaty and thereafter funded by Monies contributed by State Parties as determined by the Conference of State Parties to be paid into AMA budget.

AMA will be an advisory institution and a platform for collaboration between national medicines regulatory authorities/ RECs steered by state parties.

MIN. NO. NA/DC-H/2023/67: SUBMISSIONS BY DIFFERENT AGENCIES

a) Federation of Kenya pharmaceutical manufacturers

- Algeria, Ghana, Morocco- These countries protect their industries and local manufacturing- what will be the impact on Kenya? What is the situation with regard

to the EAC Harmonization policy? Will money be earned from registration and authorization of medicines?

- Involvement of pharmacists in research on medicines- need to have a harmonized way of training pharmacists.
- Need to ensure proper disposal of medication.
- There is fear that the pharmaceutical market might be swamped.
- Need to reduce the cost of production so as to improve productivity and cost effectiveness
- EAC working on regulation of pharmacists which will ensure access to quality products
- The Treaty is aligned to the Constitution and policy framework. It makes economic sense and strengthens regulatory expertise and ensures harmonized processes
- There is need to ensure Kenya can manufacture and benefit from the sale of Active Pharmaceutical Ingredients (APIs)?

b) Mission For Essential Drugs and Supplies (MEDS)

- MEDS supports ratification of the Treaty as it will reduce bureaucracy in manufacturing of health products and technologies due to improvement in regulatory requirements
- Kenya needs to leverage on the local capacity to ensure quality control
- There is need to do post medical surveillance
- There is need to fast-track the Kenya Development Authority Bill
- There is need to revive the Traditional Practice Bill that was done during the previous Parliament.

c) Renal Patients Society of Kenya

- The Treaty will help in reduction in the cost of drugs and ensure availability of drugs. There is need to ensure local or regional production of drugs using locally available materials and expertise so that drugs can be sold at a cheapest price. This will solve the problem of expensive drugs while ensuring safety and quality. The AMA to provide for a means of capacity building within the continent
- Africa was considered last in the supply of vaccines during the covid-19 period.

d) Kenya Association for Pharmaceutical Industry

The Treaty will ensure regulatory harmonization and convergence and tackle the following challenges

- (a) Delay in product registration which affects market authorization certification as well as product retention in the market- The EAC takes 90 to 180 days to authorize the sale of drugs within the EAC Member States.
- (b) Lengthy WHO pre-qualification processes which lengthens the provision of approvals-it takes at least four (4) years to get such pre-qualification.
- (c) Trade barriers- The Treaty incorporates best practices by providing mechanisms for inspection and audit of drugs. It also links trade and regulation by removing the multiplicity of regulation for instance Kenya has Kenyan Bureau of Standards (KEBS) and Anti-Counterfeit Authority.

e) **Drugs for Neglected Diseases**

- Accelerated access to treatment should focus on policy adoption and will there be reliance on WHO for recommendation before adoption of medicines in the continent?
- In relation to the regulatory aspect, does establishment of AMA eliminate the WHO?
- What is the effect on customs within the respective jurisdictions? Each and every jurisdiction has to inspect a drug manufacturer's factory to check quality and standards?

f) **AMREF**

- The Treaty ensures access to medicine.
- Kenya is the lead in Africa in the pharmaceutical manufacturing sector as it currently manufactures both for the local and external markets. 70% of Kenyan pharmaceutical products are exported
- The Treaty encourages capacity building through knowledge transfer and training

g) **Pharmaceutical Society of Kenya (Psk)**

- The Treaty encourages local manufacturing.
- In relation to resourcing, the Treaty is presently being funded by donors, what will happen later when donors pull out?
- The role of pharmacists and the pharmaceutical profession needs to be leveraged upon in the implementation of the Treaty.

h) **Kenya Pharmaceutical Association**

- The Association supports the treaty and memorandum by the Ministry of Foreign Affairs.
- The focus of Treaty is on governance structure. There is little on practice and the access to health care. How will HRH/healthcare workers benefit from the Treaty? Is there provision for Cross-border practice? Will there be standardization of HRH? What are the norms for HRH that will ensure harmonization of training and certification of health care professionals within the continent?
- Some countries have zero regulations. What will be the effect of the lack of domestic regulations particularly with border entry points restrictions? How does the Treaty guarantee the authenticity of health products?
- How are middle level practitioners taken care of under the Treaty?

i) **Kenya AIDS NGOs Consortium.**

- There is need to establish an African Digital Platform that lists the approved medical products and technologies in the Continent.
- The Association applauded the Kenyan Parliament for undertaking public participation on Treaty.

j) **Kenya Medical Laboratory Technicians and Technologist Board**

- The Association supports the Treaty.

k) **National Quality Control Lab**

There is need to develop an alert system for substandard and falsified medicines in the continent and the treaty should provide for the regulation of herbal products as the treaty currently only provides for dietary supplements and traditional medicine.

1) Pharmacy and Poisons Board

The Board supports the Treaty and was part of its development process and that the treaty will deal with the high cost of manufacturing of medicines and particularly APIs. South Africa and DRC have been extracting quinine. China is the biggest producer of APIs in the country. There is need to do capacity building within country and outside the country.

There is a huge market for the country as AMA is continental. Manufacturers can distribute drugs within the whole continent and manufacturers can come and set up their businesses in the country. There is therefore need to look at the cost of electricity and taxation so as to encourage the influx of pharmaceutical manufacturers hence the need to critically assess the therapeutic outcome of the treaty and tap into it as a country.

There is multiplicity of Regulations on drugs within the country. There is need to avoid duplicity for instance the country has inspectors from PPB and the Anti-Counterfeit Authority have the same responsibilities? The lack of harmonization of laws is the main reason why Kenya is at the maturity level 1. This issue to be addressed in the Kenya Drugs Authority Bill sponsored by the Chairperson of the Committee.

The Treaty will ensure short marketing timelines and will lead to revenue generation as the country can sell drugs across the continent.

There is a big challenge of capacity in the regulatory entities. The number of people in regulatory authorities is currently less than 15 in number when optimal ought to be 100. The implementation of the Treaty requires digitization of processes and systems.

The WHO prequalification process only assesses a class of drugs namely malaria, ARVs, Reproductive health, diarrhea in children, Covid-19 and other emergency issues. The Treaty will not replace the WHO but will work alongside WHO for strengthening of capacity within the country and the region. There are assessors working for WHO in Tanzania, Ghana and Kenya. Kenya has two WHO qualified assessors.

There is need to check the substandard and falsified medicines while checking market control in line with the Head of Public Service circular on border management. Cross border trading will benefit from the treaty as there is harmonized regulation. There is however need to build an alert system.

The list of OTC poisons in part I and Part II of the Schedule to the Pharmacy and Poisons Act, Cap. 244. The list needs to be revised. The law also needs to be amended to provide for scheduling and rescheduling of medicines.

There are three categories of drugs namely OTC, prescription and pharmacy only. The prescription category has more categories and the PPB has developed some guidelines on the same.

Importance of the AMA treaty

1. AMA will promote cooperation, harmonization and mutual recognition of regulatory decision in the continent.
2. AMA will reduce time taken for medication to the market and also reduce various inspections.
3. AMA will complement and enhance collaboration and contribute to improving patient's access to quality, safe and efficacious medical products and health technologies on the continent.

4. AMA will complement their efforts and contribute to their capacity building towards improving Access to quality assured medical products with the agenda of achieving universal health coverage.
5. AMA will enable Elimination of duplication through best shared practices and data exchange.
6. AMA will enable capacity building which includes knowledge transfers and trainings this will improve capacity of the states to deal with emerging issues.
7. AMA will contribute to the improved regulations of medicines, medical products and technologies and will open access to new health technologies.
8. With AMA the cost of the drugs will reduce because of the use of the local material.
9. AMA will lead to availability of drugs and vaccines this is because production will be done in Africa and every time when there are emergencies the country cannot rely on other countries.
10. The ratification of the treaty will reduce some challenges which include;
 1. Long time taken for pre-qualification by WHO.
 2. Delay of product registration and authorized certificate.

MIN. NO. NA/DC-H/2023/68: ADJOURNMENT

There being no any other business, The Chairperson, adjourned the meeting at exactly 1.00 p.m.

Sign.....Date.....

HON. DR. ROBERT PUKOSE, M.P – CHAIRPERSON

DEPARTMENTAL COMMITTEE ON HEALTH

**MINUTES OF THE TWELFTH (12TH) SITTING OF THE DEPARTMENTAL
COMMITTEE ON HEALTH HELD IN MEDIA CENTRE ON THURSDAY 16th
FEBRUARY, 2023 AT 10.30 A.M.**

PRESENT

1. The Hon. Dr. Pukose Robert, M.P -Chairperson.
2. The Hon. Ntwiga Patrick Munene, M.P -Vice-Chairperson.
3. The Hon. Sunkuli Julius Lekakeny Ole, EGH, EBS, M.P
4. The Hon. Titus Khamala, M.P
5. The Hon. Prof. Jaldesa Guyo Waqo, M.P.
6. The Hon. Kipngor Reuben Kiborek, M.P
7. The Hon. Owino Martin Peters, M.P.
8. The Hon. Wanyonyi Martin Pepela, M.P
9. The Hon. Lenguris Pauline, M.P
10. The Hon. Mary Maingi, MP.
11. The Hon. Dr. Nyikal James Wambura, M.P.
12. The Hon. Oron Joshua Odongo, M.P.
13. The Hon. Kibagendi Antony, M.P.
14. The Hon. Mathenge Duncan Maina, M.P

MEMBERS ABSENT WITH APOLOGY

1. The Hon. Muge Cynthia Jepkosgei, M.P.

COMMITTEE SECRETARIAT

- | | |
|-------------------------------|------------------------|
| 1. Mr. Hassan A. Arale | - Clerk Assistant II |
| 2. Ms. Gladys Kiprotich | - Clerk Assistant III |
| 3. Ms. Priscillah Saidi | - Research Officer III |
| 4. Ms. Faith Chepkemoi Rotich | -Legal counsel II |
| 5. Mr. Hiram Kimuhu | - Fiscal Analyst III |
| 6. Mr. Salat Abdi Ali | - Serjeant-At-Arms II |

MIN. NO. NA/DC-H/2023/48: PRELIMINARIES/INTRODUCTION

The meeting was called to order at 10.30 a.m with a word of prayer by the Hon. Dr. Pukose Robert, M.P.-Chairperson.

MIN. NO. NA/DC-H/2023/49: ADOPTION OF THE AGENDA

The agenda was adopted having been proposed by the Hon. Kibagendi Antony, M.P. and seconded by the Hon. Titus Khamala, M.P. The agenda was amended to include a brief by the Legal Counsel on the Ratification of the African Union Treaty for the Establishment of the African Medicines Agency (AMA)

MIN. NO. NA/DC-H/2023/50: BRIEFING BY THE LEGAL COUNSEL ON HEALTH (AMENDMENT BILL NO.42 OF 2022) BY HON. DISMUS BARASA, M.P, THE STATUTE LAW (MISCELLANEOUS) AMENDMENT BILL, 2022 ON MENTAL HEALTH ACT AND PHARMACY AND POISONS ACT AND RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

The Legal Counsel highlighted the content of the health (Amendment) (No. 2) Bill, No. 42 of 2022 and the comments by the stakeholders (Coalition of Blood for Africa, Kenya Healthcare Foundation and the Ministry of Health). It was noted that the Ministry of Health had requested the Committee to withhold the Bill and allow the Ministry to enforce planned interventions in ensuring an efficient and effective referral system.

After extensive deliberations, the Committee rejected the Bill for the following reasons:

- 1) Referral within the country is already provided for within the Act and is a matter of policy and needs no legislation.
- 2) Referral outside the Country is also a matter of policy that is best handled by the Ministry and does not need to be legislated upon. Legislating on the same would cause unnecessary bureaucracies to the detriment of Kenyans seeking treatment abroad.

The Committee was briefed on Statute Law (Miscellaneous) Amendment Bill, 2022 which had proposed amendments to the Mental Health Act and Pharmacy and Poisons Act by deleting the expression “Attorney-General” and replacing it with the expression “Director of Public Prosecutions) in section 42(5) and 40(4) respectively. It was noted that amendments were superfluous as these subsections (the subject of amendment) were non-existent as they had already amended in the past by Parliament by the Mental Health (Amendment) Act, No. 27 of 2022 and Health Laws (Amendment) Act, No. 5 of 2019.

The Committee thereafter resolved that, the proposed amendments to the Mental Health Act and Pharmacy and Poisons Act should be deleted from the Statute Law (Miscellaneous) Amendment Bill, 2022.

The Legal Counsel further stated that in the Memorandum of Objects and Reasons to the Statute Law (Miscellaneous) Amendment Bill, 2022, for the statutes where there is a proposed substitution of the expression “Attorney General” with the expression “Director of Public Prosecutions”, the justification for the substitutions should be indicated as ensuring harmony with the functions of the DPP under Article 157(6) and not Article 157(2) of the Constitution. The former makes provision for the functions of the DPP while the latter deals with the manner of appointment of a DPP.

Ratification of the African Union Treaty for The Establishment of The African Medicines Agency (AMA).

The Legal Counsel made a presentation on the Treaty and Memorandum submitted by the Ministry of Foreign Affairs on the Treaty. She explained the objectives and implication of the

Treaty on the country and the implementation framework for the Treaty. She also highlighted the role of Parliament in the ratification of the Treaty.

She reported that various stakeholders namely the State Department for East African Community, Ministry of East African Community and Regional Development, Ministry of Foreign and Diaspora Affairs, Office of the Attorney General, Ministry of Health, Pharmaceutical Society of Kenya (PSK), Coalition for Health Research and Development, KANCO, Kenya Revenue Authority, Office of the Attorney General and Department of Justice had expressed their support for the ratification of the Treaty.

The Chairperson reported that PATH has undertaken to host a public participation stakeholder forum for the Committee to engage the stakeholders on the Treaty on 27th February 2023.

MIN. NO. NA/DC-H/2023/51: ADOPTION OF THE REPORT ON THE SUPPLEMENTARY 1 BUDGET ESTIMATES I FOR FY 2021/2022 MINISTRY OF HEALTH

The report was adopted having been proposed by The Hon. Owino Martin Peters, M.P. and seconded The Hon. Oron Joshua Odongo, M.P.

MIN. NO. NA/DC-H/2023/52: COMMITTEE OBSERVATIONS

The Committee made the following observations during the review and scrutiny of the FY 2022/203 Supplementary Estimates:

1. That there was reduction of the total budget for State department for Medical Services vote 1081 from Kshs.122.52 billion to Kshs.66.57 billion. The recurrent budget has reduced from Kshs.68 billion to Kshs.66.57 billion and the development budget has reduced from Kshs.54.02 billion to Kshs.46.91 billion.
2. That part of the budget has been moved to the new Budget vote for the state department Public Health and Professional Standards of Kshs.5.58 billion under vote 1083. This being its first allocation both the total recurrent and development budgets have moved from zero to Kshs.2.95 billion and Kshs.2.64 billion respectively.
3. That there was reduction of the total recurrent budget by Kshs.1.93billion from Kshs.68.50billion to Kshs.66.57billion.
4. That there was reduction in the Development budget by Kshs.7.11billion from the initial allocation of Kshs.54.02billion to Kshs.46.91billion.
5. That the committee is not involved in the important events like TB world day, Malaria world day of the MoH despite its core role in over sighting the MoH.
6. That there was a reduction of the donor funds across the projects being supported/ funded by the donors.
7. The Ministry has a lot of pending bills amounting to Kshs.5billion and some of the pending bills are historical bills.
8. There was reduction of Kshs1 billion (donor funds) from the project at Cancer Centre at Kisii Level 5 but the targets as indicted in supplementary estimates is 100 %completion rate by end of the financial year 2022/23.

9. That there was no legal framework that was provided to show how the Ministry of Health can finance the county government hospitals like the Mama Margret Kenyatta hospital which is being funded by Kenyatta National Hospital.

MIN. NO. NA/DC-H/2023/53: COMMITTEE RECOMMENDATIONS

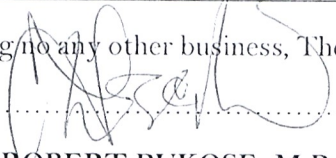
The Committee having received submissions from the Ministry of Health and having had lengthy and insightful deliberations on the proposed Supplementary Budget Estimates 1 for FY 2022/23, made the following recommendations: -

The Committee recommends the following; - That,

- 1) Proper scrutiny of all the outstanding legal claims should be conducted before authorization of any budget in future.
- 2) Supply of Medical Equipment for COVID-19-JICA grant be approved since it is a grant by Japan International Cooperation Agency (JICA).
- 3) The MoH should submit the contractual document for the MES project; this will enable the Committee to make informed decisions even in future.
- 4) Those hospitals not benefiting from MES equipment and yet they have been paying for the lease of the equipment should be facilitated so that the citizens can get the necessary services.
- 5) The Ministry of health should fully align its programmes, functions and operations as per the executive order 1 of 2023 to fit into the two state departments of the Ministry of Health.
- 6) All monies being given to level 4 hospitals should be utilized for upgrading of the same to national referral hospitals in all the 47 counties as is stated in first schedule of the health Act.

MIN. NO. NA/DC-H/2023/54: ADJOURNMENT

There being no any other business, The Chairperson, adjourned the meeting at exactly 2.00 p.m.

Sign.......... Date. 23/12/2023.....

HON. DR. ROBERT PUKOSE, M.P – CHAIRPERSON

STAKEHOLDER'S SUBMISSIONS

RATIFYING THE AFRICA MEDICINE AGENCY TREATY: A LONG JOURNEY BUT TIMELY VENTURE

The African Union (AU) Heads of State and Government adopted and endorsed the treaty for the establishment of AMA on 11 February 2019 in Addis Ababa, Ethiopia. At least 15 governments of the AU must ratify the treaty for the African Medicines Agency (AMA) to come into force. This is a unique opportunity for the African continent to tackle well-known challenges in access to quality health products and make progress towards universal health coverage.

Notably, the Constitution of Kenya in Article 43 (1) (a) **guarantees each Kenyan citizen the right to access the highest attainable standard of health including reproductive health care.** The right to health is one of the most important right for human condition without which the exercise of other rights is indispensable.

The aim of Africa's new medicine agency is to harmonize the regulatory system for medical products across the continent's 55 nations to enable faster approval processes and to support local pharmaceutical production. It is a similar model to that of the European Medicines Agency.

The primary objective of AMA is to deliver effective national regulatory frameworks with first-rate technical support on a regional and continental scale. No one nation has the resources or capacity to adequately oversee the whole supply chain for health products in today's increasingly globalized globe.

In order to increase access to essential pharmaceuticals and health commodities that are secure, efficient, economical, and of high quality, AMA will leverage African regulatory capacities and assets.

KANCO is a premier membership organization of Non-Governmental Organizations, Community Based Organizations, Faith Based-Organizations, PLHIV support organizations, learning institutions, public and private organizations responding to HIV, TB and public health Concerns. KANCO currently has a cumulative membership of 1,200 organizations and manages programs in the Eastern Africa.

Accordingly, KANCO and the Kenyan Civil Society applaud the Cabinet and the Ministry of Foreign Affairs for ratifying the AMA Treaty in May 2022. We applaud the Kenyan Parliament for requesting public memoranda, stressing the significance of key stakeholders' involvement, such as regulators, researchers, academic institutions, commercial enterprises, and civil society organizations.

To:

- Support the growth of local pharmaceutical production, a key objective of the Pharmaceutical Manufacturing Plan for Africa (PMPA)
- Provide a mechanism for evaluating medical products for the treatment of priority diseases as determined by the African Union.

- Support regularly inspects, coordinate and share information about products that are authorized for marketing.
- Coordinate joint reviews of clinical trial applications for vaccines and assessment
- Develop common standards and regulations, harmonizing legislation;
- Collaborate with RECs and National Medicines Regulatory Authorities (NMRAs) in the identification of substandard and falsified medical products (SFs) and facilitate information sharing across countries.

In maintaining and sustaining our advocacy work, as we mobilize political leaders; **We urge the Kenyan Parliament to expedite the process that will formally ratify the AMA pact by Kenya.**

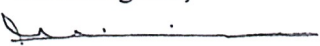
The AMA Treaty has already been ratified or signed by 33 of the 55 member states of the African Union. 20 AU nations have ratified and deposited the treaty, and three more have ratified but not yet done so. 22 countries have neither ratified nor signed, while 10 countries have signed but not ratified. Kenya belongs to the latter group.

A crucial step before the ratification of the AMA treaty can be formally lodged with the African Union is the Kenyan parliament's approval of the ratification. Kenya should lead other African nations in ratifying the pact because it is a major player in the field of global health.

The Constitution of Kenya in Article 43 (1) (a) **guarantees each Kenyan citizen the right to access the highest attainable standard of health including reproductive health care.** The right to health is one of the most important right for human condition without which the exercise of other rights is indispensable.

9TH February 2023

Kind Regards,


Allan Ragi

Executive Director

KANCO

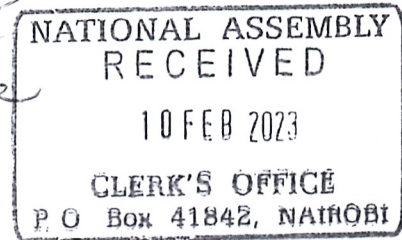


*3/ Deptal
Committee
This relates
to the new
draft
to
Health
Treaty
14-2-23*

9th February 2023

To,
Clerk of the National Assembly,
Parliament Building,
P.O Box 41842-00200,
Nairobi Kenya.

*Hassan Asale
pls facilitate
up with
15/2/23*



Dear Mr. Samuel Njoroge,

REF: SUBMISSION OF MEMORANDUM FOR CONSIDERATION BY THE NATIONAL ASSEMBLY ON THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

We the undersigned organizations under the Coalition for Health Research and Development (CHReaD) take this opportunity to urge the National Assembly to ratify Africa Medicines Agency (AMA) and enable Kenya to join hands with other progressive Africa Member states in the realization of the African Union Agenda 2063 and in making Africa a healthier and productive continent. Additionally, AMA ratification will catalyze the realization of Universal Health Coverage (UHC) by assuring faster access to the highest quality of medical products. UHC is the vehicle for government to deliver the right to health to all Kenyans as guaranteed in article 43 of the constitutional of Kenya, 2010.

CHReaD is a network that brings together Civil Society Organizations (CSOs), academia, and the private sector to advocate for an enabling policy, regulatory and financing environment that facilitates research, innovation, development of and access to medical products and technologies such as vaccines to improve the health of Kenyans.

CHReaD appreciates the government of Kenya in its effort and intention to ratify AMA. We equally appreciate the parliament of Kenya for initiating public participation on AMA ratification. We are in full support of the need to harmonize regulatory systems for medical products in Kenya and across the continent through such initiatives. AMA will provide a platform for coordination and strengthening on-going regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources.

Why should Kenya ratify AMA?

1. The African Medicines Agency is in line with the Kenya 2010 Constitution and will contribute to the achievement of the Kenya Vision 2030 while supporting the objectives of the Kenya Health Policy (2014 – 2030), the Kenya Health Sector Strategic Framework 2018–2023, and aiding the realization of the objects of the Kenya National Pharmaceutical Policy (KNPP).

2. With over 30 pharmaceutical manufacturing plants, Kenya's pharmaceutical industry is the largest in the Common Market for the Eastern and Southern Africa region (COMESA). However, Kenya still imports more than 70 percent of its pharmaceutical needs. The sector also relies heavily on imported raw materials for production. AMA will enable a conducive environment for a thriving manufacturing ecosystem that in return will spur growth of the economy and create more job opportunities for Kenyans.
3. AMA is central in ensuring the thriving and development of Africa's Pharma Industry, reducing over-reliance in imported and often expensive medicines and medical products. Kenya therefore needs to work with AMA allowing her cooperation with other AU member countries to encourage a handful of globally competitive industry clusters to thrive in country within the Regional Economic Communities (RECs) and continentally.
4. AMA will expedite Kenyans access to quality, effective and efficacious medical products due to increased local manufacture of these products that are specific to the disease burden in the country.

VALUE PROPOSITION OF RATIFICATION OF AMA TREATY

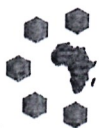
1.1 Access to complex/specialized medical products and technologies

1. AMA will provide a platform where countries such as Kenya will have access to complex, specialized and expensive molecules. With the economies of scale, the overall cost of acquiring these products will substantively reduce which eventually will result into lower product cost at the patient level.
2. AMA will guarantee access to quality specialized medical products like biologicals that have few regulatory expertise in Africa through access to rigorous pooled regulatory process.

1.2 Increase Local Manufacturing Capabilities

1. AMA will eliminate duplication and ensure efficient use of resources, towards improving access to safe and efficient medical products. AMA will provide guidance, streamline and enhance efforts of the Regional Economic Communities (RECs) and partner states (NMRAs) towards harmonization of medical products regulation. Products that Kenya exports or wishes to export shall be meeting the standards of all 55 countries in Africa. Given Kenya's advancement in manufacturing compared to many African countries, this would be a plus for its companies that do not have to try to meet different standards in different countries.
2. AMA will open up the market for Kenya's local production and manufacturing industry to the USD 1.2Billion market in Africa, which would go beyond the current USD 160Million at the EAC level.
3. AMA will create and manage an active pharmaceutical ingredients (API) database, and increase the knowledge on and reliability of APIs on the continent by capitalizing on existing API data & API inspections.

1.3 Ease of Doing Business



CHR&D

Fast-tracking health R&D, innovation, and access.

1. Increased reliance under the AMA framework will reduce the timelines for registration of medicines that would in turn reduce operational costs and pose as an incentive for manufacturers to set up factories within the region thus attracting investments.
2. AMA will increase economic integration of the continent, which will be reinforced by the African Continental Free Trade Agreement, leading to easier circulation of quality products that meet AMA and global standards, and effective control of medical products.

1.4 Capacity Building

1. AMA harmonizes technical and procedural standards across all States Parties, by coordinating guideline development and endorsement at the continent level. Harmonized technical and procedural standards (e.g., guidance) for the regulatory components along the life cycle of medical products are key enablers for successful capacity building across the continent.
2. AMA will provide capacity building to regulatory authorities such as the Pharmacy and Poisons Board (PPB) of Member states towards their regulatory maturity levels upon a request by the member states. Kenya is aiming at attaining maturity level 3 (ML3).
3. AMA as a continental agency will, through harmonization, capacity-building and dedicated activities, improve the capacity of States Parties to face the challenges of emerging and re-emerging diseases such as COVID-19 including improving of access to safe and efficacious medicines of their populations while fostering confidence and reliance among member states.

CHReaD highly recommends the expedited ratification of the AMA treaty by the Kenya government as a matter of national interest. This will put the country on the course to self-reliance in providing safe, quality and affordable medical products as well as significantly improve our economic status.

Signed:

1. AMREF HEALTH AFRICA IN KENYA
2. PATH
3. IAVI
4. KENYA AIDS NGOs CONSORTIUM (KANCO)
5. MOVEMENT OF MEN AGAINST AIDS IN KENYA (MMAAK)
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8. Q_DATAMS
9. CONCERN WORLDWIDE
10. NATIONAL ORGANIZATION OF PEER EDUCATORS (NOPE)
11. VILLGRO AFRICA
12. INTERNATIONAL COUNCIL OF WOMEN FIGHTING AIDS IN KENYA (ICWK)

13. NATIONAL EMPOWERMENT NETWORK OF PEOPLE LIVING WITH HIV IN KENYA (NEPHAK)
14. ACCESS TO MEDICINES PLATFORM
15. AMBASSADOR FOR YOUTH AND ADOLESCENT REPRODUCTIVE HEALTH PROGRAM (AYARHEP)
16. KENYA TREATMENT ACCESS MOVEMENT (KETAM)
17. DEUTSCHE STIFTUNG WELTBEVÖLKERUNG (DSW) KENYA
18. MÉDECINS SANS FRONTIÈRES (MSF)
19. KENYA PROGRESSIVE NURSES ASSOCIATION.
20. CENTRAL KENYA DEVELOPMENT NETWORK



REPUBLIC OF KENYA

OFFICE OF THE ATTORNEY-GENERAL
&
DEPARTMENT OF JUSTICE

*D/Dept Law
Committees*

Your Ref: NA/DCS/DC-H/2022/008
Our Ref: AG/CONF/6/E/85 VOL XIII

2nd February 2023

Mr. Samuel Njoroge
The Clerk
National Assembly
NAIROBI

*Hassan Abalo
pls facilitate
4/2/23
9/2/23*

RE: CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA) BY THE GOVERNMENT OF KENYA

Reference is made to the above captioned subject matter.

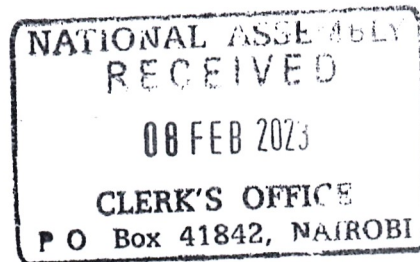
We hereby acknowledge receipt of your letter referenced NA/DCS/DC-H/2022/008 and dated 25th January 2023, the contents of which are duly noted.

We have reviewed the full text of the above referenced Treaty and the accompanying Memoranda to Parliament and find the same to be in order from a legal perspective.

Thank you for your continued cooperation.

KENNEDY OGETO CBS
SOLICITOR GENERAL

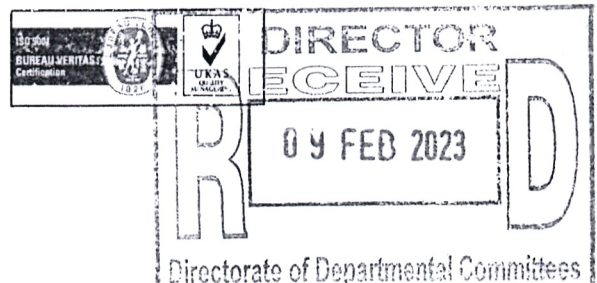
Copy to: Hon. J.B.N Muturi, EGH
ATTORNEY GENERAL



SHERIA HOUSE, HARAMBEE AVENUE
P.O. Box 40112-00100, NAIROBI, KENYA. TEL: +254 20 2227461/2251355/07119445555/0732529995
E-MAIL: info.state.law.office@kenya.go.ke WEBSITE: www.attorney-general.go.ke

DEPARTMENT OF JUSTICE
CO-OPERATIVE BANK HOUSE, HAILLE SELLAHIE AVENUE P.O. Box 56057-00200, Nairobi-Kenya TEL: Nairobi 2224029/ 2240337
E-MAIL: legal@justice.go.ke WEBSITE: www.justice.go.ke

ISO 9001:2008 Certified



*3/Departmental Committees
Is the Treaty before the committee? explain process of ratification. IF not, the process of ratification and participation in. 9/2/23*

*Hassan Asale
pls TNA
13/2/23*

9th February 2023

To,
Clerk of the National Assembly,
Parliament Building, P.O Box 41842-00200,
Nairobi Kenya.

NATIONAL ASSEMBLY
RECEIVED
09 FEB 2023
CLERK'S OFFICE
P. O. Box 41842, NAIROBI

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DIRECTOR
RECEIVED
13 FEB 2023
P.O. Box 30125-00100, Nairobi, Kenya | www.amref.org/kenya/chread/
Directorate of Departmental Committees

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12. INTERNATIONAL COUNCIL OF WOMEN FIGHTING AIDS IN KENYA (ICWK)



CHR_{ea}D

Fast-tracking health R&D, innovation, and access.

13. NATIONAL EMPOWERMENT NETWORK OF PEOPLE LIVING WITH HIV IN KENYA (NEPHAK)
 14. ACCESS TO MEDICINES PLATFORM
 15. AMBASSADOR FOR YOUTH AND ADOLESCENT REPRODUCTIVE HEALTH PROGRAM (AYARHEP)
 16. KENYA TREATMENT ACCESS MOVEMENT (KETAM)
 17. DEUTSCHE STIFTUNG WELTBEVÖLKERUNG (DSW) KENYA
 18. MÉDECINS SANS FRONTIÈRES (MSF)
 19. KENYA PROGRESSIVE NURSES ASSOCIATION.
 20. CENTRAL KENYA DEVELOPMENT NETWORK
- CC: Clerk of the National Assembly- Health Committee**

SECRET



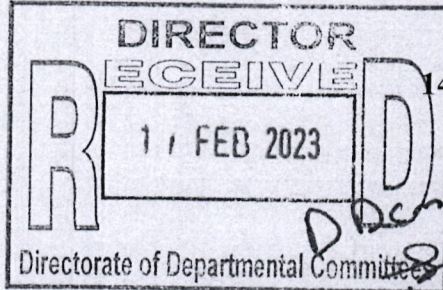
**REPUBLIC OF KENYA
THE NATIONAL TREASURY AND ECONOMIC PLANNING**

Telegraphic Address: 22921
FINANCE – NAIROBI
Fax No. 315779
Telephone: 2252299
When replying please quote

THE NATIONAL TREASURY
P.O. Box 30007 - 00100
NAIROBI
KENYA

Ref: TNT/CONF 70/02 TY⁷ (50)

Samwel Njoroge
Clerk of the National Assembly
Parliament Buildings
NAIROBI



14th February, 2023

Dear

Mr Njoroge

Hassan Arabe 16/2/23
to facilitate 17/2/23

RE: MEMORANDUM BY THE CABINET SECRETARY FOR THE NATIONAL TREASURY AND ECONOMIC PLANNING ON THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY BY THE GOVERNMENT OF KENYA

Reference is made to your letter Ref. No. NA/DCS/DC-H/2022/008 dated 24th January, 2023 on the subject matter.

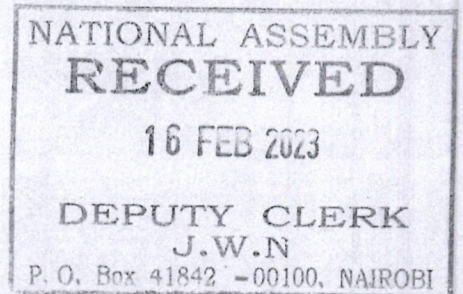
We wish to note that the National Treasury and Economic Planning has no objection to the ratification of the Treaty as noted in the enclosed Memorandum by the Cabinet Secretary for the National Treasury and Economic Planning

The purpose of this letter therefore is to forward the Memorandum for your necessary action.

Yours

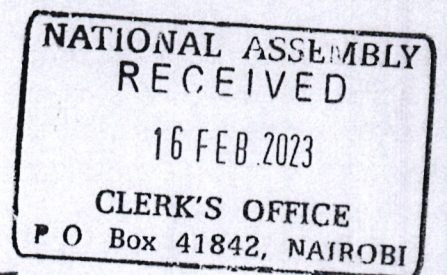
Sincerely
[Signature]

DR. CHRIS K. KIPTOO, CBS
PRINCIPAL SECRETARY/THE NATIONAL TREASURY



Copy to:

Prof. Njuguna Ndung'u, CBS
Cabinet Secretary
National Treasury and Economic Planning
NAIROBI



SECRET



FEDERATION OF KENYA PHARMACEUTICAL MANUFACTURERS

P.O. Box 53362-00200 NAIROBI. KENYA

Panesar Center, Mombasa Rd after Airtel, Opposite Auto Express, 2nd Floor, Room 206

Tel: +254 723916980

E-mail: info@fkpm.co.ke Website; www.fkpm.co.ke

Our Ref: FKPM/160/03/2023/NA

1st March 2023

To,

Mr. Samuel Njoroge
Clerk of the National Assembly,
Parliament Buildings,
Nairobi.

2/Dep'tal Comms
Hassan Arabe
FK Facilitator
W 1/10
5/3/23
"Advance copy by email"

Dear Sir,

REF: FKPM COMMENTS ON THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE AFRICAN MEDICINES AGENCY (AMA).

Please receive warm greetings from FKPM (Federation of Kenyan Pharmaceutical Manufacturers) which is an organization representing the interests of Local Pharmaceutical Manufacturers by conveying them to the relevant forums regarding policies, regulations, trade and quality.

We write this in reference to the recent Memorandum on the ratification of the African Union Treaty for the establishment of the African Medicines Agency (AMA).

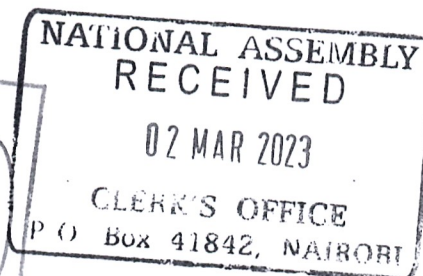
As the Federation of the Local Pharmaceutical Manufacturers, we support the ratification process, however we have some comments we would like to bring to your attention:

1. There are some protected countries, i.e. Algeria, Morocco (Support sale of locally manufactured products) and Ghana (Have a restricted list of products for which they don't allow importation) that were mentioned and we would like to understand how other African Companies can penetrate these markets so that it's a fair playing ground for all.
2. We would like to further understand what mechanisms have been put in place to ensure that the African Medicines Agency (AMA) will work with the Individual State parties and Regional Economic Communities to regulate Medical Products in the African Region.
3. We would like to request that Kenya as a country should be represented in the Secretariat of the African Medicines Agency (AMA).

We look forward to you favourable response

Yours sincerely,


Dr. Vimal Patel
Chairman



*D/D Dept
Commisar.*

NATIONAL ASSEMBLY
RECEIVED
07 MAR 2023
CLERK'S OFFICE
P. O. Box 41842, NAIROBI



LAB MEDICINE SOCIETY OF KENYA
LUNGA LUNGA SQUARE, 3RD FLOOR, UNIT 332, LUNGA LUNGA ROAD
P. O. BOX 35992-00200, NAIROBI.
TEL.: +254737320000 EMAIL: labmedicinekenya@gmail.com

Our Ref. No: LMSK/02/23/Vol.2

Date: 6TH/03/2023

CLERK OF THE NATIONAL ASSEMBLY
REPUBLIC OF KENYA,
P.O BOX 41842-00100,
NAIROBI.

*Hasan Asale
fks faalidate
w w w
8/3/23*

SUBJECT: MEMORANDUM ON THE RATIFICATION OF AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF AFRICAN MEDICINE AGENCY.

Introduction

Lab Medicine Society of Kenya (LMSK) is a professional society whose membership comprises of Medical Laboratory Officers and Scientists in Kenya practicing in hospitals laboratories and medical research institutions.

Rationale

In strengthening and harmonization of regulation of medical products in Africa Union member states and Regional Economic Communities (RECs), LMSK just like other institutions whose members use medical products in offering essential healthcare services to patients has analyzed the above mentioned treaty, understood its provisions and hereby gives its proposals understood to be appropriate for effectiveness of objectives of the treaty, effectiveness of regulatory policies that shall arise from the treaty, quality & safety of medical products that shall be in market and effectiveness of application of those medical products.

We therefore propose the following amendment:

DIRECTOR
RECEIVED
08 MAR 2023
Directorate of Departmental Committees

1. That, the name “**African Medicines Agency**” is not the correct name to be used in the yet to be formed agency of African Union and should be amended to “**African Medical Products Agency**”. We cite the below justification to get your understanding of this our proposal.
 - a. Refer to the definition of the term “**Medical Product**” and the term “**Medicine**” under **page 6 of ARTICLE 2**. “Medical product” in the definition covers a wide list of items such as medicines, vaccines, blood, blood products and diagnostic devices & medical devices. “Medicines” in the definition narrows down to substance or a mixture of substances. This means that “medicines” as an item is just one among examples of medical products.

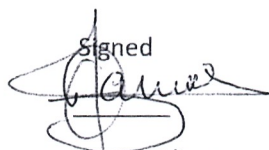
Further, the term medicine as defined excludes medical diagnostic devices, donated blood for transfusion and donated tissues for transplant which currently are and shall continue to be examples of medical products under regulation in every member state of AU. Considering that “Medical diagnostic devices”, include laboratory in vitro testing kits, laboratory machines used in analysis of human specimens, radiology machines among other machines which in real sense are not “substances or mixture of substances” hence singly cannot be categorized as medicines.

We therefore understand that the term **AFRICAN MEDICAL PRODUCTS AGENCY** is the best name which includes all preparations under healthcare use.
 - b. Refer to main objective of African Medicines Agency under **ARTICLE 4 page 8** of the document; it states: To enhance capacity of state parties and Regional Economic Communities to regulate **medical products** in order to improve access to quality, safe and efficacious **medical products** on the continent. The conspicuous, repetitive and appropriate term here is **medical products**.
 - c. Refer to **article 6 page 10**, under functions of the Agency. The key word as evidenced in functions; a, b, d, e and f is “Medical Products” (which is an inclusive list of healthcare items and preparations under use) and not “Medicines”.
2. We support the treaty having realized its positive impacts on the continent of Africa.

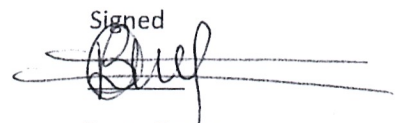
Thank you.

Signed


Phelix Otieno
President (LMSK)

Signed


Barasa Oliver
Secretary General (LMSK)

Signed


Bruce Onyango
Treasurer (LMSK)

27th February, 2023

The Clerk of the National Assembly,

P.O. Box 41842-00100,

NAIROBI.

Mr Araki
Please deal
03/03/2023

RE: MEMORANDA BY KENYA MEDICAL LABORATORY TECHNICIANS & TECHNOLOGISTS BOARD (KMLTTB) ON THE MATTER OF CONSIDERATION BY NATIONAL ASSEMBLY ON THE RATIFICATION OF THE AFRICAN UNION TREATY ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

27TH FEBRUARY 2023, MERCURE HOTEL, UPPER HILL (Formerly Crowne Plaza).

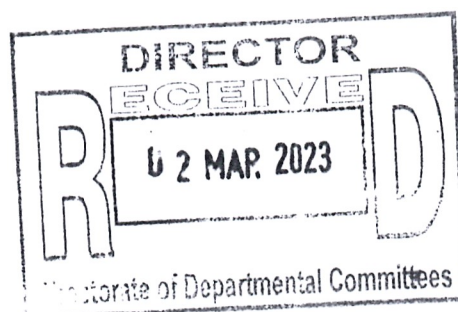
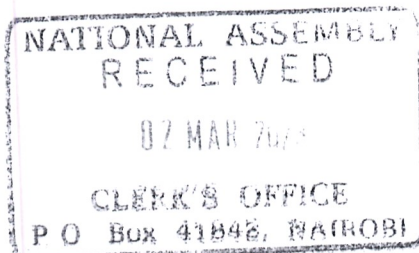
The Kenya Medical Laboratory Technicians and Technologists Board (KMLTTB) is a body corporate with the mandate to exercise general supervision and control over the training, business, practice and employment of Laboratory Technicians and Technologists in Kenya and to advise the Government in relation to all aspects thereof as provided in CAP 253A and legal notice No.113 of 2011.

KMLTTB Supports the ratification of the AMA as it will go a long way in bringing positive impact to Kenya and other member states of the African union. The Agency will indeed go a long way in ensuring access to safe and quality medicines in Africa.

However, we would like to make some distinction between drugs intended in the treaty and Invitro Diagnostic Devices (IVDs) also known as invitro diagnostic health products.

KMLTTB would wish to inform parliament of Kenya to appreciate the difference between the medicines/drugs and medical devices/health products and technologies envisaged by the Treaty against Invitro diagnostic health products.

IVDs which have a high potential risk on the population have a precise performance criteria officially listed in the common technical specifications, which are the standards that control IVDs (*European Union MDR 2017*).



The fundamental differences between Medicines /drugs, health products & technologies and IVDs lies in the following aspects; -

Medicines/drugs, health products & technologies	Invitro Diagnostic Products
Pharmaceutical products of therapeutic and preventive purposes.	Medical Laboratory reagents, kits, controls, calibrators, software, instruments and systems) intended for use in the in diagnosis of disease or other conditions, including a determination of the state of health in order to cure, mitigate, treat or prevent diseases. Such products are intended for use in the collection, preparation, and examination of specimens taken from the human body (<i>Iso 15189:2022</i>) (<i>Iso 13485</i>).
They come into contact with the patient	They do not come in direct contact with the patient
They have a claim of therapeutic effect	No therapeutic effect claimed
They fulfill their role based on direct action on the patient.	They fulfill their role based on information they provide and not on direct action on the patient. Instead the quality of information delivered by an IVD is assessed by measuring the analytical precision of the test or assay and by the clinical evidence of the information provided

Whereas it is important to seek a high level of safety for both medical devices & IVDs, it is equally desirable to maintain the difference between these two categories of medical devices – one that comes in direct contact and the other that does not, to ensure a rational and effective regulatory system.

KMLTTB wishes that the Kenyan Parliament and indeed the Parliaments of all member states maintain the quality of Invitro diagnostic products. This will have positive impact on the objectives of AMA. Also help avert misdiagnosis and mismanagement of disease conditions as well as strengthen the mandates of the respective regulatory authorities.

Respectfully,



Patrick Kisabei - Ag REGISTRAR – KMLTTB

Cc. The Hansard National Assembly.

Cc. Parliamentary Committee on Health.

27th February, 2023

Mr Amole
Desl
03/03/23

The Clerk of the National Assembly,

P.O. Box 41842-00100,

NAIROBI.

RE: MEMORANDA BY KENYA MEDICAL LABORATORY TECHNICIANS & TECHNOLOGISTS BOARD (KMLTTB) ON THE MATTER OF CONSIDERATION BY NATIONAL ASSEMBLY ON THE RATIFICATION OF THE AFRICAN UNION TREATY ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

27TH FEBRUARY 2023, MERCURE HOTEL, UPPER HILL (Formerly Crowne Plaza).

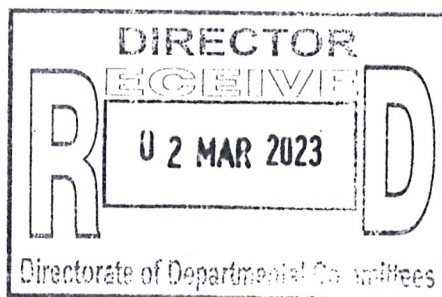
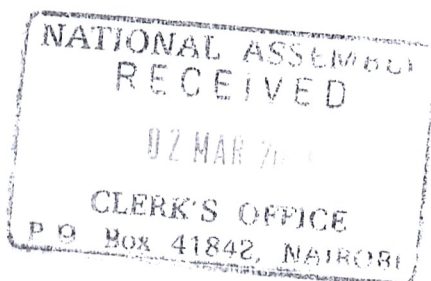
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Respectfully,



Patrick Kisabei - Ag REGISTRAR – KMLTTB

Cc. The Hansard National Assembly.

Cc. Parliamentary Committee on Health.

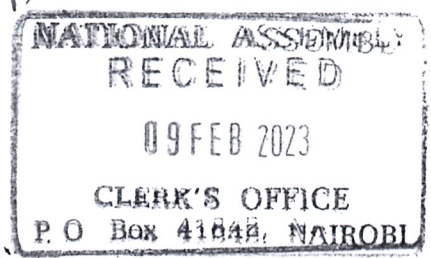
*DLS
To respond, receipt
and explain
to the
the process;
the shared
the signing.
Executive
CS
9/2/23*



PHARMACEUTICAL SOCIETY OF KENYA
 PCEA Foundation, Block C, Jabavu Rd.
 P.O. Box 44290 - 00100 Nairobi - KENYA
 T. +254 722 817 264
 E. info@psk.or.ke
 W. www.psk.or.ke

8th February 2023

Mr. Samuel Njoroge,
 Clerk of the National Assembly,
 P.O. Box 41842-00100,
 Nairobi,



Dear Mr. Njoroge,

Re: The request dated 26th January 2023 requesting public participation memoranda on the treaty to establish the African Medicines Agency

The Pharmaceutical Society of Kenya (PSK) is the professional body of Pharmacists in Kenya, mandated to advance the practice of Pharmacy, to advocate for and to promote the welfare of its members. Our goal is to advance ethical practice of pharmacy in Kenya and to advocate for better pharmaceutical policies in the health sector for the benefit of Kenyans. PSK recognizes that pharmacists are uniquely and ideally positioned in communities to provide an expanded set of promotive, preventive, and clinical health services to expand the reach of the health system to communities, families and individuals.

Pharmacists are trained to provide Effective, Efficient, Safe, Quality, Affordable and Appropriate pharmaceutical care (the practice and philosophy of appropriate medication therapy use). The achievement of the UHC (Universal Healthcare Coverage) SDG#3 (Sustainable Development Goals) goal requires resource stewardship and optimization of therapies. Right drug for the Right condition. As per FIP (International Pharmaceutical Federation) research, For every 1 (Kshs) paid to a pharmacist, the healthcare system saves 12 (Kshs).

The first 2 pages of the African Medicine Agency treaty clearly states the current situation. Subsequently and recognizes that many organizations are working to improve the situation. As PSK, we propose the following.

- 1) That the Pharmacist be proritized in the discovery and innovation of new therapeutic products locally. Being the medication experts, Pharmacist are uniquely educated and have the intellectual and technical skill to support the innovation of new therapeutic agents. This is currently underutilized and underfunded.
- 2) That the pharmacists ensure that new therapies are well designed for safety and efficacy and the that the end product is a quality product

NATIONAL EXECUTIVES: DR. LOUIS S. MACHOGU (PRESIDENT) | DR. ANGELINE ACHOKA (NATIONAL TREASURER)
 DR. DANIELLA MUNENE (V.P. GOVERNANCE) | DR. JOSEPH KATHARE (V.P. PRACTICE) | DR. TABITHA KIMANI (V.P. LOBBYING)

NEW FACE OF PHARMACY: Public | Academia | Trade, Commerce & Enterprise | Epidemiology | Hospital & Specialisation
 Leadership & Admin | Manufacturing | Community | Clinical Research | Nutraceuticals | Regulatory & Quality Assurance
 Medical Waste Management | Herbal Medicine | Health & Wellness Coach | Systems & Processes | Supply Chain | Agrovet
 Statistics & Analytics | Pharmaco Economics | Post Market Surveillance | e-Health | Drug Discovery | Sports Medicine
 Green Cross | Equipment | Clinical Pharmacy | Human Resources for Pharmacy | Practice Standards | Policy & Legislation

BEST PRACTICE | PUBLIC INTEREST | PROFESSIONALISM



PHARMACEUTICAL SOCIETY OF KENYA

PCEA Foundation, Block C, Jabavu Rd.
P.O. Box 44290 - 00100 Nairobi - KENYA

T. +254 722 817 264

E. info@psk.or.ke

W. www.psk.or.ke

- 3) That the pharmacist be involved with the process of choosing the appropriate therapeutic product for the right patient, right condition, right strength, right dose, and right frequency. The active involvement of pharmacists in pharmaceutical therapy decision is a critical part of healthcare optimization.
- 4) That the pharmacist be responsible for the safety and monitoring of the patient's therapy.
- 5) That the pharmacist be responsible for the procurement and handling of therapeutic products.
- 6) That the pharmacist be responsible for disposal of unused and expired medications in a manner that is safe to the public and environment.
- 7) That the pharmacist will access, recommend and contribute to patient safety regarding therapy choice, medication interaction and organ function, will require that all inpatient facilities have a ratio of pharmacist to in patient bed.
- 8) That the pharmacist counsel all patients inpatient and outpatient before pharmaceutical care starts.
- 9) That the pharmacist have the knowledge and skills to respond to drug toxicity challenges
- 10) That the pharmacist must be involved in public/ population health.
- 11) That the pharmacist leads the development, education and maintenance of drug formulary. Pharmacist will collect data and use it in advising on drug formulary matters.
- 12) That the pharmacist will develop, educate on, and maintain a document that will guide the levels of practice in the pharmacy profession.
- 13) That the pharmacist will form a think tank that will be funded by AU and or AMA to collect data, and offer policy direction and guidance on the effective use of medicines. The think tank will be tasked with responding to the changing needs of pharmacy.
- 14) That the pharmacist will work with higher learning regulators to develop a harmonized curriculum that produce pharmacists with the skills to match the current needs and is flexible enough to accommodate future needs. The curriculum should reflect an African consensus and to produce a practice ready pharmacist that can work anywhere in Africa.
- 15) That the pharmacist will be the driver of drug systems technologies guided by need best access current and future patients and healthcare system needs.

A USA study on economic cost of medication error concluded that, '... the economic cost of medication errors is US\$ 20-40 Billion a year...' if extrapolated, the economic cost of medication errors in Kenya is upwards of between US\$3- 6 Billion. The combined cost of providing healthcare in Kenya is approximately US\$6 Billion. Does it mean that appropriate deployment of the pharmacists could enable Kenya meet the UHC painlessly?

NATIONAL EXECUTIVES: DR. LOUIS S. MACHOGU (PRESIDENT) | DR. ANGELINE ACHOKA (NATIONAL TREASURER)
DR. DANIELLA MUNENE (V.P. GOVERNANCE) | DR. JOSEPH KATHARE (V.P. PRACTICE) | DR. TABITHA KIMANI (V.P. LOBBYING)

NEW FACE OF PHARMACY: Public | Academia | Trade, Commerce & Enterprise | Epidemiology | Hospital & Specialisation
Leadership & Admin | Manufacturing | Community | Clinical Research | Nutraceuticals | Regulatory & Quality Assurance
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BEST PRACTICE | PUBLIC INTEREST | PROFESSIONALISM



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The Treaty For The Establishment of the African Medicines Agency, clear objectives regarding medicines are to; ensure public safety, promote innovation, streamline product registration, harmonize healthcare systems, harmonize healthcare education systems, ensure positive healthcare outcomes for patients, and optimize healthcare expenditures. It is clear from the above outlined roles that a pharmacist is an important stakeholder and should be included in the entire process of establishing the proposed agency.

Africa is moving towards Non-Communicable-Diseases where patients will need daily dose of medications in addition to life style to have quality of life and enjoy relationships. Pharmacists are medication experts. By involving pharmacist in all aspect of NCD Care the patient will get optimal pharmaceutical care, positive healthcare outcomes and ultimately the slowing down of disease progression. This will reduce cost of healthcare provision and lead to the UHC that we all strive to achieve.

It is with great confidence that we believe that you will accept and act on our proposal for the sake of the patients, their families, their friends, and our economy. We all have a role to play to build our economies to make our countries and continent better than we found them. Pharmacists can help by keeping you safe and healthy.

Sincerely,

Dr. Louis. S. Machogu
President,
Pharmaceutical Society of Kenya

NATIONAL EXECUTIVES: DR. LOUIS S. **MACHOGU** (PRESIDENT) | DR. ANGELINE **ACHOKA** (NATIONAL TREASURER)
DR. DANIELLA **MUNENE** (V.P. GOVERNANCE) | DR. JOSEPH **KATHARE** (V.P. PRACTICE) | DR. TABITHA **KIMANI** (V.P. LOBBYING)

NEW FACE OF PHARMACY: Public | Academia | Trade, Commerce & Enterprise | Epidemiology | Hospital & Specialisation
Leadership & Admin | Manufacturing | Community | Clinical Research | Nutraceuticals | Regulatory & Quality Assurance
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Statistics & Analytics | Pharmaco Economics | Post Market Surveillance | e-Health | Drug Discovery | Sports Medicine
Green Cross | Equipment | Clinical Pharmacy | Human Resources for Pharmacy | Practice Standards | Policy & Legislation

BEST PRACTICE | PUBLIC INTEREST | PROFESSIONALISM

KANCO

Healthy people, empowered communities

Nkoro, Merisho
Past O/Rongai Off Magadi Road, Acacia-Matasia Road,
Plainsview North Road, Opposite Merisho Police Post
P.O. Box 69866 – 00400
Nairobi, Kenya

Tel: +254 20 2323506 / 2323533
+254 20 2434615 / 2322657
Cell phone: +254 722 203 344
Email: kanco@kanco.org
Website: www.kanco.org

February 9, 2023

To,

Mr. Samuel Njoroge
Clerk of the National Assembly
P.O. Box 41842-00100
Nairobi

Dear Sir,


REF: SUBMISSION OF MEMORANDUM ON THE RATIFICATION OF THE ESTABLISHMENT OF THE AFRICA MEDICAL AGENCY TREATY.

I refer to the above subject matter.

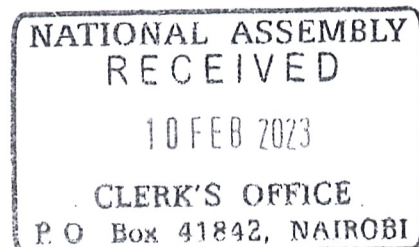
Please find attached the memorandum.

All assistance will be highly appreciated.

Kind Regards,


Allan Ragi
Executive Director
KANCO

*DHS DEPT
70
advise them of
process
the memo
take
request
and
interest
C/S
under
per.
10/2/23*



RATIFYING THE AFRICA MEDICINE AGENCY TREATY: A LONG JOURNEY BUT TIMELY VENTURE

The African Union (AU) Heads of State and Government adopted and endorsed the treaty for the establishment of AMA on 11 February 2019 in Addis Ababa, Ethiopia. At least 15 governments of the AU must ratify the treaty for the African Medicines Agency (AMA) to come into force. This is a unique opportunity for the African continent to tackle well-known challenges in access to quality health products and make progress towards universal health coverage.

Notably, the Constitution of Kenya in Article 43 (1) (a) **guarantees each Kenyan citizen the right to access the highest attainable standard of health including reproductive health care.** The right to health is one of the most important right for human condition without which the exercise of other rights is indispensable.

The aim of Africa's new medicine agency is to harmonize the regulatory system for medical products across the continent's 55 nations to enable faster approval processes and to support local pharmaceutical production. It is a similar model to that of the European Medicines Agency.

The primary objective of AMA is to deliver effective national regulatory frameworks with first-rate technical support on a regional and continental scale. No one nation has the resources or capacity to adequately oversee the whole supply chain for health products in today's increasingly globalized globe.

In order to increase access to essential pharmaceuticals and health commodities that are secure, efficient, economical, and of high quality, AMA will leverage African regulatory capacities and assets.

KANCO is a premier membership organization of Non-Governmental Organizations, Community Based Organizations, Faith Based-Organizations, PLHIV support organizations, learning institutions, public and private organizations responding to HIV, TB and public health Concerns. KANCO currently has a cumulative membership of 1,200 organizations and manages programs in the Eastern Africa.

Accordingly, KANCO and the Kenyan Civil Society applaud the Cabinet and the Ministry of Foreign Affairs for ratifying the AMA Treaty in May 2022. We applaud the Kenyan Parliament for requesting public memoranda, stressing the significance of key stakeholders' involvement, such as regulators, researchers, academic institutions, commercial enterprises, and civil society organizations.

To:

- Support the growth of local pharmaceutical production, a key objective of the Pharmaceutical Manufacturing Plan for Africa (PMPA)
- Provide a mechanism for evaluating medical products for the treatment of priority diseases as determined by the African Union.

- Support regularly inspects, coordinate and share information about products that are authorized for marketing.
- Coordinate joint reviews of clinical trial applications for vaccines and assessment
- Develop common standards and regulations, harmonizing legislation;
- Collaborate with RECs and National Medicines Regulatory Authorities (NMRAs) in the identification of substandard and falsified medical products (SFs) and facilitate information sharing across countries.

In maintaining and sustaining our advocacy work, as we mobilize political leaders; **We urge the Kenyan Parliament to expedite the process that will formally ratify the AMA pact by Kenya.**

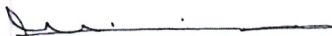
The AMA Treaty has already been ratified or signed by 33 of the 55 member states of the African Union. 20 AU nations have ratified and deposited the treaty, and three more have ratified but not yet done so. 22 countries have neither ratified nor signed, while 10 countries have signed but not ratified. Kenya belongs to the latter group.

A crucial step before the ratification of the AMA treaty can be formally lodged with the African Union is the Kenyan parliament's approval of the ratification. Kenya should lead other African nations in ratifying the pact because it is a major player in the field of global health.

The Constitution of Kenya in Article 43 (1) (a) **guarantees each Kenyan citizen the right to access the highest attainable standard of health including reproductive health care.** The right to health is one of the most important right for human condition without which the exercise of other rights is indispensable.

9TH February 2023

Kind Regards,



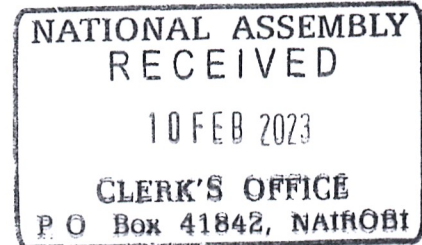
Allan Ragi

Executive Director

KANCO

9th February 2023

To,
Clerk of the National Assembly,
Parliament Building,
P.O Box 41842-00200,
Nairobi Kenya.



Dear Mr. Samuel Njoroge,

REF: SUBMISSION OF MEMORANDUM FOR CONSIDERATION BY THE NATIONAL ASSEMBLY ON THE RATIFICATION OF THE AFRICAN UNION TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

We the undersigned organizations under the Coalition for Health Research and Development (CHReaD) take this opportunity to urge the National Assembly to ratify Africa Medicines Agency (AMA) and enable Kenya to join hands with other progressive Africa Member states in the realization of the African Union Agenda 2063 and in making Africa a healthier and productive continent. Additionally, AMA ratification will catalyze the realization of Universal Health Coverage (UHC) by assuring faster access to the highest quality of medical products. UHC is the vehicle for government to deliver the right to health to all Kenyans as guaranteed in article 43 of the constitutional of Kenya, 2010.

CHReaD is a network that brings together Civil Society Organizations (CSOs), academia, and the private sector to advocate for an enabling policy, regulatory and financing environment that facilitates research, innovation, development of and access to medical products and technologies such as vaccines to improve the health of Kenyans.

CHReaD appreciates the government of Kenya in its effort and intention to ratify AMA. We equally appreciate the parliament of Kenya for initiating public participation on AMA ratification. We are in full support of the need to harmonize regulatory systems for medical products in Kenya and across the continent through such initiatives. AMA will provide a platform for coordination and strengthening on-going regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources.

Why should Kenya ratify AMA?

1. The African Medicines Agency is in line with the Kenya 2010 Constitution and will contribute to the achievement of the Kenya Vision 2030 while supporting the objectives of the Kenya Health Policy (2014 – 2030), the Kenya Health Sector Strategic Framework 2018–2023, and aiding the realization of the objects of the Kenya National Pharmaceutical Policy (KNPP).

2. With over 30 pharmaceutical manufacturing plants, Kenya's pharmaceutical industry is the largest in the Common Market for the Eastern and Southern Africa region (COMESA). However, Kenya still imports more than 70 percent of its pharmaceutical needs. The sector also relies heavily on imported raw materials for production. AMA will enable a conducive environment for a thriving manufacturing ecosystem that in return will spur growth of the economy and create more job opportunities for Kenyans.
3. AMA is central in ensuring the thriving and development of Africa's Pharma Industry, reducing over-reliance in imported and often expensive medicines and medical products. Kenya therefore needs to work with AMA allowing her cooperation with other AU member countries to encourage a handful of globally competitive industry clusters to thrive in country within the Regional Economic Communities (RECs) and continentally.
4. AMA will expedite Kenyans access to quality, effective and efficacious medical products due to increased local manufacture of these products that are specific to the disease burden in the country.

VALUE PROPOSITION OF RATIFICATION OF AMA TREATY

1.1 Access to complex/specialized medical products and technologies

1. AMA will provide a platform where countries such as Kenya will have access to complex, specialized and expensive molecules. With the economies of scale, the overall cost of acquiring these products will substantively reduce which eventually will result into lower product cost at the patient level.
2. AMA will guarantee access to quality specialized medical products like biologicals that have few regulatory expertise in Africa through access to rigorous pooled regulatory process.

1.2 Increase Local Manufacturing Capabilities

1. AMA will eliminate duplication and ensure efficient use of resources, towards improving access to safe and efficient medical products. AMA will provide guidance, streamline and enhance efforts of the Regional Economic Communities (RECs) and partner states (NMRAs) towards harmonization of medical products regulation. Products that Kenya exports or wishes to export shall be meeting the standards of all 55 countries in Africa. Given Kenya's advancement in manufacturing compared to many African countries, this would be a plus for its companies that do not have to try to meet different standards in different countries.
2. AMA will open up the market for Kenya's local production and manufacturing industry to the USD 1.2Billion market in Africa, which would go beyond the current USD 160Million at the EAC level.
3. AMA will create and manage an active pharmaceutical ingredients (API) database, and increase the knowledge on and reliability of APIs on the continent by capitalizing on existing API data & API inspections.

1.3 Ease of Doing Business



1. Increased reliance under the AMA framework will reduce the timelines for registration of medicines that would in turn reduce operational costs and pose as an incentive for manufacturers to set up factories within the region thus attracting investments.
2. AMA will increase economic integration of the continent, which will be reinforced by the African Continental Free Trade Agreement, leading to easier circulation of quality products that meet AMA and global standards, and effective control of medical products.

1.4 Capacity Building

1. AMA harmonizes technical and procedural standards across all States Parties, by coordinating guideline development and endorsement at the continent level. Harmonized technical and procedural standards (e.g., guidance) for the regulatory components along the life cycle of medical products are key enablers for successful capacity building across the continent.
2. AMA will provide capacity building to regulatory authorities such as the Pharmacy and Poisons Board (PPB) of Member states towards their regulatory maturity levels upon a request by the member states. Kenya is aiming at attaining maturity level 3 (ML3).
3. AMA as a continental agency will, through harmonization, capacity-building and dedicated activities, improve the capacity of States Parties to face the challenges of emerging and re-emerging diseases such as COVID-19 including improving of access to safe and efficacious medicines of their populations while fostering confidence and reliance among member states.

CHReaD highly recommends the expedited ratification of the AMA treaty by the Kenya government as a matter of national interest. This will put the country on the course to self-reliance in providing safe, quality and affordable medical products as well as significantly improve our economic status.

Signed:

1. AMREF HEALTH AFRICA IN KENYA
2. PATH
3. IAVI
4. KENYA AIDS NGOs CONSORTIUM (KANCO)
5. MOVEMENT OF MEN AGAINST AIDS IN KENYA (MMAAK)
6. WACI HEALTH
7. STOP TB PARTNERSHIP KENYA
8. Q_DATAMS
9. CONCERN WORLDWIDE
10. NATIONAL ORGANIZATION OF PEER EDUCATORS (NOPE)
11. VILGRO AFRICA
12. INTERNATIONAL COUNCIL OF WOMEN FIGHTING AIDS IN KENYA (ICWK)



CHR^{ea}D

Fast-tracking health R&D, innovation, and access.

13. NATIONAL EMPOWERMENT NETWORK OF PEOPLE LIVING WITH HIV IN KENYA (NEPHAK)
14. ACCESS TO MEDICINES PLATFORM
15. AMBASSADOR FOR YOUTH AND ADOLESCENT REPRODUCTIVE HEALTH PROGRAM (AYARHEP)
16. KENYA TREATMENT ACCESS MOVEMENT (KETAM)
17. DEUTSCHE STIFTUNG WELTBEVÖLKERUNG (DSW) KENYA
18. MÉDECINS SANS FRONTIÈRES (MSF)
19. KENYA PROGRESSIVE NURSES ASSOCIATION.
20. CENTRAL KENYA DEVELOPMENT NETWORK

Telephone: +254-20-318888
Fax: +254-20-2240066/341935/344333
Email: communication@mfa.go.ke
Website: www.mfa.go.ke
When replying please quote Ref. No. and date



HARAMBEE AVENUE
P.O. Box 30551-00100
NAIROBI, KENYA

MINISTRY OF FOREIGN AFFAIRS

MFA/RT/TCA./11/VOL. I

3rd February, 2023

Ref. No.....

Mr. Samuel Njoroge
Clerk of the National Assembly
Parliament Buildings
P.O Box 41842-00100
NAIROBI

Dear Sir,

*3/ Dept of the Hon. Secy
To process
Hassan Arala
pls facilitate
7/2/23*

**RE: CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION
TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES
AGENCY (AMA) BY THE GOVERNMENT OF KENYA**

Reference is made to the above subject matter and your letter dated 25th January, 2023 referenced NA/DCS/DC-H/2022/008.

According to the Treaty Making and Ratification Act, the Ministry of Foreign & Diaspora Affairs is mandated to manage Kenya's International Agreements and Commitments including all procedures related to the mechanisms of initiation, signature, ratification, implementation, reporting of compliance and review of Agreements.

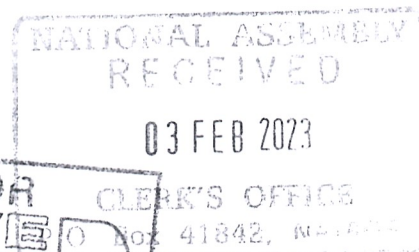
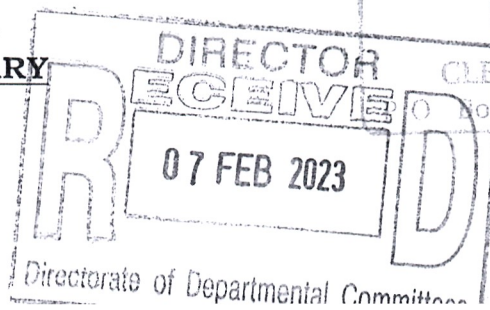
The Ministry of Foreign & Diaspora Affairs convened a stakeholder meeting with the Office of the Attorney General, and the Ministry of Health nominated representatives and came up with a joint brief on the AMA memoranda.

The purpose of this letter is to, therefore, submit the joint brief for your consideration.

Yours

Sincerely

**Dr. A. Korir Sing'Oei,
PRINCIPAL SECRETARY**





**KENYA REVENUE
AUTHORITY**

ISO 9001:2015 CERTIFIED

Office of the Commissioner General

KRA/5/1002/5(8502)

13th February 2023

Mr. Samuel Njoroge
Clerk of the National Assembly
P. O. Box 41842-00100
Parliament Buildings
Nairobi

Dear *C Clerk,*

*DCS
20
PROCESS
14/2/23*

*Hassan Arale
to facilitate
15/2/23*

**CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION
TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES
AGENCY (AMA) BY THE GOVERNMENT OF KENYA**

Reference is made to your letter Ref: NA/DCS/DC-H/2022/008 dated 25th January 2023 on the above subject matter.

The Authority has reviewed the Memorandum on the Ratification of the *African Treaty for the Establishment of the African Medicines Agency (AMA)* together with the Treaty for the Establishment of the African Medicines Agency and does not have additional input on the document for submission.

Yours sincerely,

**FCPA Githii Mburu, MGH, CBS
COMMISSIONER GENERAL**

**DIRECTOR
RECEIVED
15 FEB 2023
Directorate of Departmental Committees**

**NATIONAL ASSEMBLY
RECEIVED
14 FEB 2023
CLERK'S OFFICE
P O Box 41842, NAIROBI**



REPUBLIC OF KENYA

NATIONAL ASSEMBLY
RECEIVED
03 FEB 2023
CLERK'S OFFICE
P.O. Box 41842, NAIROBI

**MINISTRY OF EAST AFRICAN COMMUNITY AND REGIONAL
DEVELOPMENT**

STATE DEPARTMENT FOR EAST AFRICAN COMMUNITY

Telephone: +254-20-2245741/2211614/2245752
Mobile: 0729111108/0733208888
Wireless: +254-20 2603599/20 2603733
E-mail: ps@meac.go.ke
Website: www.meac.go.ke
When replying please quote:

Co-op Bank House Building
Haile Selassie Avenue
P.O. Box 8846-00200 City Square
NAIROBI, KENYA

*Dept of
Communities*

EAC/5/8/Vol. 50(114)

2nd FEBRUARY, 2022

The Clerk,
National Assembly,
P.O Box 41842 – 00100,
NAIROBI.
Email: cna@parliament.go.ke

*Hassan Arabe
pls facilitate
w m w
7/2/23*

**RE: CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION TREATY
FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)
BY THE GOVERNMENT OF KENYA**

Reference is made to your letter, Ref. NA/DCS/DC-H/2022/008 dated 25th January, 2023 on the above captioned subject.

The State Department for East African Community has considered the provisions of the Treaty for the Establishment of the Africa Medicines Agency (AMA) and consulted with the Ministry of Health and is of the view that Kenya should consider ratifying the Treaty.

In view of this, please find attached our memoranda in support of ratification of the treaty for your consideration.

Odibe

**Charity Chepkonga-Bokindo
FOR: PRINCIPAL SECRETARY**

Encl.

**DIRECTOR
RECEIVED
07 FEB 2023**
Directorate of Departmental Committees



Vision:
*Deepen and widen East African Integration for Sustainable Development and improved livelihoods of all
Kenyans*



MEMORANDA ON RATIFICATION OF THE TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

JUSTIFICATION FOR RATIFICATION

1. Since 2009, the Africa Union Development Agency (AUDA-NEPAD), working with regional economic communities (RECs) and collaborating with development partners, has been advancing the African Medicines Regulatory Harmonization (AMRH) Initiative that has now culminated into the African Medicines Agency (AMA).
2. The AMA is intended to provide a platform for coordination and strengthening on-going regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources. This memorandum therefore seeks to support Kenya's ratification.
3. AMA will enable Kenya strengthen its Clinical Trials ecosystem including that of COVID-19, strengthen its manufacturing industry, enable it conform to the best practices and standard for health products, strengthen Kenya's capacity to regulate and monitor safety of health products.
4. AMA will provide a platform for a multi-faceted approach for combating Substandard and Falsified medical products by strengthening medicine regulatory systems including the capacity for conducting pre-marketing authorizations and routine post marketing surveillance.
5. The existing national and regional regulatory bodies or harmonization initiatives at RECs level will continue with their mandate but AMA will complement their efforts and contribute to capacity building towards improving access to quality-assured medical products within the agenda of Universal Health Coverage and the Sustainable Development Goals.
6. Kenya supported the AMRH initiatives including in the set-up of the EAC-MRH Programme and implementation. These initiatives at the regional and continental levels have largely aided Kenya's realization of its Health Sector development goals and targets while strengthening the national capacities for effective health service delivery.
7. The establishment of AMA puts Kenya at a better standpoint to benefit more once AMA comes into force. It is therefore in the national interest that Kenya ratifies the Treaty for the Establishment of AMA



REPUBLIC OF KENYA

NATIONAL ASSEMBLY
RECEIVED
06 FEB 2023
CLERK'S OFFICE
P.O. Box 41842, NAIROBI

**MINISTRY OF EAST AFRICAN COMMUNITY AND REGIONAL
DEVELOPMENT**

STATE DEPARTMENT FOR EAST AFRICAN COMMUNITY

Telephone: +254-20-2245741/2211614/2245752
Mobile: 0729111108/0733208888
Wireless: +254-20 2603599/20 2603733
E-mail: ps@meac.go.ke
Website: www.meac.go.ke
When replying please quote:

Co-op Bank House Building
Haile Selassie Avenue
P.O. Box 8846-00200 City Square
NAIROBI, KENYA

*3/ Depted Committee
20 accounts & receipts
and register
pro call
W.M.*

EAC/5/8/Vol. 50(114)

2nd FEBRUARY, 2022

The Clerk,
National Assembly,
P.O Box 41842 – 00100,
NAIROBI.
Email: cna@parliament.go.ke

*Hassan Arale
pls factcheck
up W.M.
7/2/23*

**RE: CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION TREATY
FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)
BY THE GOVERNMENT OF KENYA**

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In view of this, please find attached our memoranda in support of ratification of the treaty for your consideration.

Atibe

**Charity Chepkonga-Bokindo
FOR: PRINCIPAL SECRETARY**

Encl.

**DIRECTOR
RECEIVED**
07 FEB 2023
Directorate of Departmental Committees



Vision:
Deepen and widen East African Integration for Sustainable Development and improved livelihoods of all
Kenyans



NEWSPAPER ADVERTISEMENT ON PUBLIC PARTICIPATION

Report The 2022 Ibrahim Index reveals the country continues to deteriorate on security and rule of law Kenya ranked 13th in governance index

The country has improved from position 18 in Africa in the 2020 report by the Mo Ibrahim Foundation

MERCY CHELANGAT

Kenya has made strides in overall governance, according to a new Africa report. The report, however, reveals the country has deteriorated on security and rule of law, inclusion and equality, foundations for economic opportunity and human development. According to the 2022 Ibrahim Index of African Governance, Kenya now ranks at position 13 on the continent, an improvement from position 18 in the 2020 report by the Mo Ibrahim Foundation. The ranking is among the 35 countries that have recorded small improvements in governance. The country still needs additional measures to address human trafficking, labour, crime levels, impartiality of judicial system, accessibility of public records, anti-corruption mechanisms and public procurement procedures. Kenya also has its work cut out ensuring that public administration is streamlined to allow for



Mo Ibrahim, the founder of Mo Ibrahim Foundation, chats with Rwanda President Paul Kagame during the Mo Ibrahim Governance Weekend in Kigali in April 2018. FILE IN NATION

AT A GLANCE

How countries performed

The five countries with the highest improvement include Seychelles, Tunisia, Botswana, Mauritius, and Cape Verde, with the latter two deteriorating over the decade. Some 22 countries, however, decreased their score since 2017 and include South Sudan, Central African Republic. Even though Somalia, Eritrea and Equatorial Guinea were ranked with the lowest score, they have improved their score over the decade.

better civil registration, improving capacity of the statistical system and mobilisation of tax and revenue collection.

According to the report, the country also needs to do better in regional integration, improve access to banking services, land and water, rural market and ensure

equal rural representation and participation.

Overall, the report warns that Africa's decade of progress may go down the drain due to failure of improvement.

The report, which is released after every two years, allows for citizens, governments and institutions to assess delivery of public goods and services and policy outcomes in Africa, details the latest available data of the African continent as from 2012 to 2021, focusing on security and rule of law, participation, rights and inclusion, foundations for economic opportunity and human development.

It details that security and rule of law in Africa deteriorated almost four times between 2019 and 2021 compared to between 2012 and 2019.


Security and accountability, and transparency have also declined at a fast pace since 2019.

Participation, rights and inclusion have also deteriorated six times faster between 2019 and 2021 as compared to between 2012 and 2019.

"The most deteriorated indicators over the last three years are Freedom of Association and Assembly and Democratic Elections. While the pace of decline has accelerated for Freedom of Association and Assembly, the deterioration in Democratic Elections reverts the progress made between 2012 and 2019," reads the report.


Even though the overall governance in Africa has improved since 2012, thanks to advances in human development and economic foundation, progress has been slow since 2019, partly due to worsening security and weaker democracy, unemployment, inefficient transport and poor energy infrastructure. Covid-19 also played a part in worsening "pre-existing deteriorations in security and democracy."

Out of the 54 African countries, 35 countries improved their governance since 2012, with only 15 accelerating their progress between 2017 and 2021. The five countries with the highest improvement include Seychelles, Tunisia, Botswana, Mauritius, and Cape Verde, with the latter two deteriorating over the decade.



THIKA TECHNICAL TRAINING INSTITUTE

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E-MAIL: thikatechnical@gmail.com, thikatech@yahoo.com
Website: www.thikatechnical.ac.ke



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
S/no	Vacancy No	Position	Desired Qualification	No. Of Position
1	TTTI/BoG/MET/01/01/2023	Mechanical Engineering Technician	Diploma in Mechanical Engineering Plant or Mechanical Engineering Refrigeration & Air Conditioning.	1
2	TTTI/BoG/ACC/02/01/2023	Accountant	<ul style="list-style-type: none"> • Bachelors of Commerce in Finance/ Accounting or its equivalent • CPA-K Certificate • Member of ICPAK 	1

NB: For more details visit the Institute website: www.thikatechnical.ac.ke

Application letter, detailed CV, photocopies of relevant certificates and testimonials should reach the Institute on or before **03/02/2023**. Clearly addressed to: -

The Principal
Thika Technical Training Institute,
P.O Box 91-01000,
Thika.

Only shortlisted candidate will be contacted.



REPUBLIC OF KENYA
THE NATIONAL ASSEMBLY
13TH PARLIAMENT – FIRST SESSION
DEPARTMENTAL COMMITTEE ON HEALTH

In the Matter of Articles 2(5) and (6) and 118 (1)(b) of the Constitution of Kenya and Section 8 of the Treaty Making and Ratification Act, 2012 and
In the Matter of Consideration by the National Assembly of the Ratification of the African Union Treaty for the Establishment of the African Medicines Agency (AMA)

**INVITATION FOR PUBLIC PARTICIPATION
(SUBMISSION OF MEMORANDA)**

Article 118(1) (b) of the Constitution of Kenya requires Parliament to facilitate public participation and involvement in the legislative and other business of Parliament and its Committees.

Pursuant to Section 8 of the Treaty Making and Ratification Act, 2012, the Cabinet Secretary for Foreign Affairs submitted to the National Assembly the *African Union Treaty for the Establishment of the African Union Medical Agency (AMA)* for approval, after it was ratified by the Cabinet on 12th May, 2022.

The Treaty seeks to establish the African Medical Agency as a specialized Agency of the African Union with its own rules, membership and resources to enhance the capacity of state parties and regional Economic Communities (RECs) to regulate medical products in order to improve access to quality, safe and sufficient medical products on the continent.

The Treaty was initially tabled in the National Assembly towards the end of the 12th Parliament and was thereafter not considered. Consequently, it was re-tabled in the House on 1st December 2022 in the 13th Parliament and thereafter committed to the Departmental Committee on Health for consideration and reporting as required under the Treaty Making and Ratification Act, 2012.

In compliance with Article 118 (1) (b) of the Constitution, the Departmental Committee on Health hereby invites the public and stakeholders to submit memoranda on the Treaty. The full text of the Treaty and the accompanying Memorandum to the National Assembly may be accessed through <http://www.parliament.go.ke/the-national-assembly/house-business/paper-laid>.

The memoranda should be addressed to the Clerk of the National Assembly, P.O. Box 41842-00100, Nairobi, hand-delivered to the Office of the Clerk of the National Assembly, Main Parliament Building, Nairobi; or emailed to cn@parliament.go.ke; to be received on or before 9th February, 2023 at 5.00 p.m.

SAMUEL NJOROGE
CLERK OF THE NATIONAL ASSEMBLY
26th January, 2023

"For the Welfare of Society and the just Government of the People"

Directorate of Legislative &
Procedural Services

The Table Office

MEMO

TO : DIRECTOR, DEPARTMENTAL COMMITTEES
FROM : AG. DEPUTY DIRECTOR, LEGISLATIVE &
PROCEDURAL SERVICES
DATE : JANUARY 18, 2023
SUBJECT : PAPER LAID

Hassan Arala
to table before the
committee for consideration
17/1/23

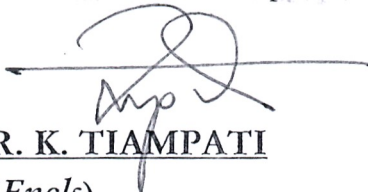
The following Paper was laid on the Table of the House on Tuesday, June 7, 2022:

- Memorandum on the Ratification of African Union Treaty for the establishment of the African Medicines Agency (AMA).

Considering that the House proceeded to *Sine die* recess immediately thereafter, marking the end of the Term of the 12th Parliament, the paper could not be considered.

The said Treaty was re-tabled before the House on Thursday, December 1, 2022 in the 13th Parliament.

Enclosed herewith, please find the said Treaty for your necessary action.


R. K. TIAMPATI
(Encls)

Copy: Clerk of the National Assembly
Deputy Clerks
Director, Legislative & Procedural Services





Head, Take this
to register, cause
taking and
signature
committee
signature
2/6/22

OFFICE OF THE CABINET SECRETARY

MFA.RT/CAB/VOL.1

18th May, 2022

Mr. Michael R. Sialai, EBS
Clerk of the National Assembly
Parliament Buildings
NAIROBI

07 JUN 2022

Tuesday (3)

DL & P
Lam

H TO
2/6/22

Dear Mr. Sialai,

Maamah Mw 2/6/22

**RE: MEMORANDUM ON THE RATIFICATION OF AFRICAN UNION
TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES
AGENCY (AMA)**

I refer to the above matter.

The Cabinet, during a Meeting held on 12th May, 2022, approved the Ratification of the African Union Treaty for the Establishment of the African Medicines Agency (AMA).

Pursuant to Section 8 of the Treaty Making and Ratification Act, 2012, I hereby submit a Memorandum and a copy of the Treaty for consideration by the National Assembly.

Yours sincerely

**AMB. RACHELLE OMAMO, SC, EGH
CABINET SECRETARY**

Encls.

02 JUN 2022

Copy to: **Dr. Joseph K. Kinnya, EGH**
Head of Public Service
Executive Office of the President
NAIROBI

Hon. Mutahi Kagwe, EGH

Cabinet Secretary

Ministry of Health

NAIROBI

Hon. (Amb.) Ukur Yatani, EGH

Cabinet Secretary

National Treasury & Planning

NAIROBI

Hon. Justice (Rtd.) P. Kihara Kariuki, EGH

Attorney General

Office of the Attorney General

& Department of Justice

Sheria House

NAIROBI





THE NATIONAL ASSEMBLY

MINISTRY OF FOREIGN AFFAIRS JUN 2022

PASSED BY:

Lom

CLEARANCE AVAILABLE:

Masonah MOW

MEMORANDUM

ON THE

RATIFICATION OF AFRICAN UNION TREATY

FOR THE ESTABLISHMENT OF THE

AFRICAN MEDICINES AGENCY (AMA)

**MEMORANDUM ON THE RATIFICATION OF AFRICAN UNION TREATY
FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY
(AMA)**

1.0 OBJECTIVE OF THE MEMORANDUM

- 1.1 The objective of this Memorandum is to seek approval for Kenya's ratification of the Ratification of African Union Treaty for the Establishment of the African Medicines Agency (AMA)
- 1.2 The ratification process was approved by the Cabinet during its meeting held on **12th May, 2022.**

2.0 BACKGROUND

- 2.1 Africa's public and private sector actors are increasingly recognizing that real region-wide progress and transformation is only attainable through improved connectivity, competitive logistics and production value chain integration in targeted strategic sectors including pharmaceuticals and agriculture. This, together with the establishment of regulatory policy convergence, is vital for the continent's trade and regional integration agenda.
- 2.2 The pharmaceutical sector, under the guidance of the African Union, has developed and launched initiatives under the Pharmaceutical Manufacturing Plan for Africa (PMPA) framework of the AU endorsed by the Assembly in 2005.
- 2.3 In 2019, due to the fact that, weak regulatory systems have resulted in the circulation of substandard and falsified (SF) medical products in many African Union Member States; posing risk to public health, harming patients and undermining confidence in healthcare delivery systems; the Assembly of Heads of State and Government, at its 32nd

ordinary session decision Assembly/AU/Dec.735(XXXII) reaffirmed the Executive Council 34th ordinary session decision EX.CL/1141(XXXIV) to establish the African Medicines Agency placing an emphasis on investment in regulatory capacity strengthening.

- 2.5 The Treaty seeks to establish the African Medicines Agency (AMA) to enhance the capacity of State Parties and Regional Economic Communities (RECs) to regulate medical products in order to improve access to quality, safe and efficacious medical products on the continent.
- 2.6 Twenty-six (26) member states (Algeria, Benin, Burundi, Cameroon, Chad, Cote d'Ivoire, Egypt, Gabon, Ghana, Guinea, Madagascar, Mali, Mauritius, Morocco, Niger, Rwanda, Republic of Congo, Saharawi Arab Democratic Republic, Senegal, Seychelles, Sierra Leone, Tanzania, Togo, Tunisia, Uganda and Zimbabwe) have signed the treaty
- 2.7 Seventeen (17) member states (Algeria, Benin, Burkina Faso, Cameroon, Chad, Gabon, Ghana, Guinea, Mali, Mauritius, Namibia, Niger, Rwanda, Seychelles, Sierra Leone, Tunisia and Zimbabwe) have ratified the Treaty for the Establishment of the African Medicines Agency and deposited the legal instrument of ratification to the Commission.
- 2.8 The Treaty for the Establishment of the African Medicines Agency (AMA) entered into force on **5th November 2021**.

3.0 OBJECT AND SUBJECT MATTER OF THE CONVENTION

- 3.1 The African Medicines Agency is a Specialized Agency of the African Union with its own rules, membership and resources to enhance the capacity of State Parties and Regional Economic Communities (RECs), to regulate medical products in order to improve access to quality, safe and efficacious medical products on the continent.

- 3.2 The AMA intends to provide a platform for coordination and strengthening of on-going regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources. The AMA will not replace existing national and regional regulatory bodies or harmonization initiatives at RECs level.
- 3.3 AMA will complement their efforts and contribute to their capacity building towards improving access to quality assured medical products within the agenda of Universal Health Coverage and the Sustainable Development Goals.
- 3.4 AMA defines acceptable standards in the regulation of medical products in the continent. The establishment of a Continental Agency that contributes to the improved regulation of medicines, medical products and technologies is therefore timely and critical.

4.0 OBLIGATIONS IMPOSED BY THE PROTOCOL

- 4.1 The AMA's vision is to ensure that all Africans have access to quality-assured, safe, efficacious and affordable medical products, that meet internationally recognised standards, for priority diseases or conditions
- 4.2 The obligations of the AMA Treaty are forward looking. States parties are obligated to *inter alia*:-
- a. To coordinate national and sub-regional medicines regulatory systems;
 - b. To conduct regulatory oversight of selected medical products including traditional medicines;
 - c. To promote cooperation, harmonisation and the mutual recognition of regulatory decision;
 - d. To strength and harmonize efforts of the African Union-recognized RECs, RHOs and Member States; and

- e. To complement and enhance collaboration and contribute to improving patients' access to quality, safe and efficacious medical products and health technologies on the continent.

5.0 PROBLEM ANALYSIS

- 5.1 An assessment performed by the World Health Organisation (WHO) of 26 African National Medicines Regulatory Authorities (NMRAs) between 2002 and 2009 found that only 15% of these NMRAs were mandated to carry out functions of marketing authorization, licensing, inspection, quality control and pharmacovigilance of medical products.
- 5.2 In many cases, not all of these functions were operational, including having access to a functional national regulatory quality control laboratory. It is important to note that not all NMRAs are expected to perform all the regulatory functions on their own, but could rely on other NMRAs' decisions such as for Good Manufacturing Practice inspection of foreign manufacturing sites and marketing authorisations.
- 5.3 The assessment found that even where local manufacturing occurred, good distribution practices (GDP) were poorly enforced thereby increasing the risk of substandard, spurious, falsely labelled, falsified and counterfeit medical products in the market. In addition to the above matters, common challenges within the continental regulatory space include lack of published standards and operating procedures, and shortage of qualified personnel.
- 5.4 Although the assessment was based on information obtained over twelve years ago, for the most part, this reflection is still valid in the current pharmaceutical regulation situation in the continent.
- 5.5 Access to quality health products and technologies, especially for low- and middle-income countries, during the COVID-19 pandemic continues to be a challenge due to disruptions in the global supply

chain systems. If established in the coming years, AMA will help African nations to fight pandemics and support national and regional responses by ensuring that only high-quality drugs, vaccines, and other health-related supplies reach African populations.

6.0 JUSTIFICATION FOR RATIFICATION

6.1 The signing and ratification of the Treaty by Kenya will demonstrate Kenya's commitment to the Continents' collective action to the improved regulation of medicines, medical products and technologies. Ratification will bring about positive consequences both to the country and States Members which include:

i) Ease of Doing Business

The AMA will provide guidance, streamline and enhance efforts of the RECs towards harmonization of medical products regulation. This will ensure efficient resource utilization by reducing duplication of investments by Member states and expediting introduction of registered medicines in the regional market through harmonized procedures.

The AMA will lead to reduction of operational costs as Kenya will employ mutual recognition of the regulatory decisions of other countries; thereby offering incentives for manufacturers to set up factories within the region thus attracting investments.

ii) Access to Safe, Quality and Efficacious Medical Products

The AMA will serve as a catalyst for stronger regulatory oversight to counteract proliferation of Substandard and Falsified medical products and enable competitiveness of locally produced medicines, particularly those used to treat diseases and conditions disproportionately affecting the African continent. This will be achieved through cross-border enforcement based on

enhanced collaboration amongst stakeholders including customs, police and judiciary.

iii) Capacity Building

Expertise built as a result of interaction by professionals from the various countries, and routine assessment of regulatory systems encourages regulatory agencies to improve processes. This translates to enhanced quality assurance systems for medical products.

iv) Access to the African Continental Free Trade Area

The African Continental Free Trade Area (AfCFTA), makes Africa the largest geographically integrated trading area in the world, allowing access without tariffs to a market of over 1.2 billion potential consumers and by extension creating an African Economic Community by 2028. This therefore will have significant implications to public health and safety, hence regulation of health products and technologies shall be critical to guaranteeing the protection of this market from fake, substandard, and counterfeit products and services.

v) African Industrialisation

The progressive industrialization of Africa, and the possibility of transforming raw materials into products, including into medicines, medical devices and technologies; requires Kenya to strategically position herself as a leader under the AU recognized RECs. This will be in line to advance the implementation of the Pharmaceutical Manufacturing Plan of Africa.

7.0 CONSTITUTIONAL AND LEGISLATIVE IMPLICATIONS

7.1 The Convention is consistent with the Constitution and promotes constitutional values and objectives, it does not allude to an amendment of the Constitution.

7.2 The Treaty advocates for the adoption of the African Union Model Law on Regulation of Medical products. This will require Kenya to amend existing relevant legislation and policies to adapt to this model law in

the spirit of harmonization to enable implementation of the Convention. Some of which may include, the Pharmacy and Poisons Act, Cap 244 of the Laws of Kenya, Health Products and Technologies Bill.

- 7.3 Kenya may also need to generate guidelines for the periodic reporting obligations generated from joint assessment exercises to establish the capacity of member states in health products and technologies and technical capacities, in line with the proposed logical framework for AMA.
- 7.4 Other non-legislative, yet practical measures that Kenya may need to undertake include: the review of existing policies and develop regionally cohesive protocols to enable participation in harmonization activities.

8.0 IMPLICATIONS RELATING TO COUNTIES

- 8.1 The obligations imposed under the Protocols are under the purview of the National Government.

9.0 FINANCIAL IMPLICATIONS

- 9.1 At the onset, the AMA will be supported by donor funding. Thereafter, States Parties will be required to contribute the amounts to be assessed by the Conference of States Parties towards the AMA budget upon the lapse of donor support funding.
- 9.2 The financial requirements during implementation will be catered for during the normal budgetary estimates of the relevant Ministries, Departments and Agencies.

10. MINISTERIAL RESPONSIBILITY

- 10.1 The Ministry that will be responsible for the implementation and any activity in regard to the Convention is the Ministry of Health.

10.2 The Office of the Attorney General and Department of Justice and the Ministry of Foreign Affairs will coordinate the reporting process on State obligations pursuant to the Treaty Making and Ratification Act No 45 of 2012.

11. RESERVATIONS

11.1 Article 35, permits member States to submit reservations when ratifying the Treaty on condition that it is compatible with the objects and purpose of the Treaty. Presently, the Ministry of Health has no reservations.

12. PUBLIC PARTICIPATION

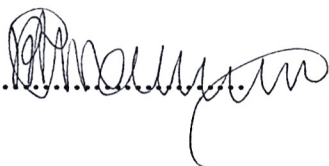
12.1 Public participation has been undertaken via various fora including and virtual meetings.

13. RECOMMENDATION TO THE NATIONAL ASSEMBLY

13.1 In consideration of the aforementioned facts, the National Assembly is invited to:

1. Note the contents of the Memorandum;
2. Consider and approve Kenya's Ratification of African Union Treaty for the Establishment of the African Medicines Agency; and
3. Direct the Cabinet Secretary of Foreign Affairs to prepare and deposit the relevant instruments to the Depository, the Chairperson of the African Union Commission.

SIGNED.....

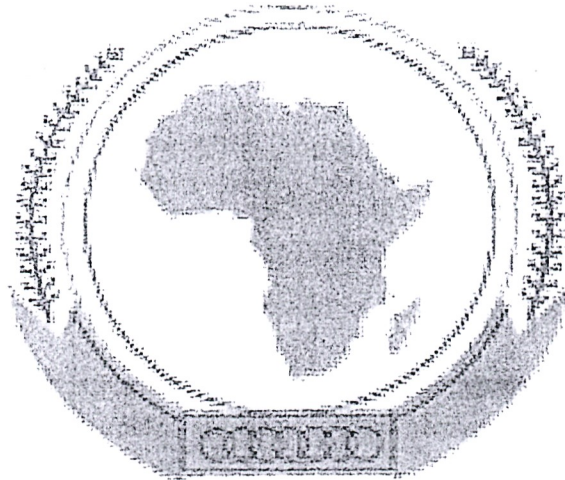


DATED.....

30th

MAY, 2022

**AMB. RAYCHELLE OMAMO, SC, EGH
CABINET SECRETARY
MINISTRY OF FOREIGN AFFAIRS**



**TREATY FOR THE ESTABLISHMENT
OF THE
AFRICAN MEDICINES AGENCY**

TREATY OF THE AFRICAN MEDICINES AGENCY (AMA)

We, Member States of the African Union,

AFFIRMING THAT quality-assured, safe and efficacious medical products are fundamental to the health and safety of the population of Africa;

AWARE THAT, weak regulatory systems have resulted in the circulation of substandard and falsified (SF) medical products in many of the African Union Member States;

COGNIZANT THAT the existence of SF products poses a risk to public health, harms patients and undermines confidence in healthcare delivery systems;

RECALLING the 55th Decision of the African Union (AU) {Assembly /AU/Dec.55 (IV)} taken during the Abuja Summit in January 2005, which requested the AU Commission to develop a Pharmaceutical Manufacturing Plan for Africa (PMPA) within the framework of the New Partnership for Africa's Development (NEPAD), aimed to improve access to good quality, safe and efficacious medical products and health technologies for the African population;

FURTHER RECALLING the Eighteenth Ordinary Session of the Heads of State and Government Orientation Committee 29 – 30 January 2012 Decision {Assembly/AU/DEC-413(XVIII)} Para 6 which endorsed the African Medicines Regulatory Harmonization (AMRH) Programme implemented through the regional economic communities (RECs);

RECOGNIZING the aspirations of the AU Roadmap on Shared Responsibility and Global Solidarity for the AIDS, tuberculosis and malaria response in Africa {Assembly AU/Dec.442 (XIX)}, Pillar II on access to medicines which aims to accelerate and strengthen regional medicines regulatory harmonization initiatives and lay the foundation for a single African regulatory agency;

BEING COGNIZANT of the challenges posed by the lack of availability of medicines and vaccines during public health emergencies of international concern and, in particular,

during the recent outbreak of the Ebola virus disease (EVD) in Africa and the attendant dearth of medical product candidates for clinical trials;

RECOGNIZING the contribution of the African Vaccines Regulatory Forum (AVAREF) in facilitating approval of EVD candidate therapies and vaccines and efforts undertaken by the African Union (AU), regional economic communities (RECs) and regional health organizations (RHOs) to mobilize human, financial and material resources and continental expertise to deal with the outbreak of EVD; and subsequent establishment of regional expert working groups (EWGs) on clinical trials oversight in East African Community (EAC) and the Economic Community of West African States (ECOWAS) as part of the implementation of the decision of the Assembly of the Union, Assembly/AU/Dec.553(XXIV) on Ebola Virus Disease (EVD) Outbreak, of January 2015);

DESIRING the use of continental institutional, scientific and regulatory resources to improve access to safe, efficacious and quality medicines; and **AWARE OF** the establishment of the African Medicines Regulatory Harmonization (AMRH) in 2009, under the management and guidance of the NEPAD Agency working with RECs and RHOs, to facilitate harmonization of regulatory requirements and practice among the national medicines regulatory authorities (NMRAs) of the AU Member States to meet internationally acceptable standards, and provide a favourable regulatory environment for pharmaceutical research and development, local production and trade across countries on the African continent;

APPRECIATING the launch and subsequent implementation of Medicines Regulatory Harmonization (MRH) Programmes and collaborative efforts in and between the East African Community (EAC); Economic Community of West African States (ECOWAS) and the West African Economic and Monetary Union (WAEMU); and the Southern African Development Community (SADC);

RECOGNIZING other on-going efforts on cooperation between the Economic Community of Central African States (ECCAS) and the Organization for Coordination in the Fight against Endemic Diseases in Central Africa (OCEAC) on implementation of the AMRH Programme in the Central African region; and the North-Eastern Africa regional

collaboration and harmonization under the leadership of the Intergovernmental Authority on Development (IGAD);

NOTING the commitment made by the African Ministers of Health during their First meeting held on 17 April 2014 in Luanda, Angola, jointly organized by the African Union Commission and World Health Organisation (WHO) to prioritize investment in regulatory capacity development; to pursue efforts towards convergence and harmonization of medical products regulation in RECs; to allocate adequate resources for the establishment of the African Medicines Agency (AMA), and the subsequent endorsement of the establishment of the AMA Task Team to spearhead the process;

RECALLING the July 2012 AU Assembly Declaration, Assembly/AU/Decl.2(XIX) on the report of AIDS Watch Africa (AWA) Action Committee of Heads of State and Government in which the Council decided that the African Medicine Regulatory Harmonization (AMRH) Initiative shall serve as a foundation for the establishment of AMA.

FURTHER RECALLING the AU Assembly Decision, Assembly/AU/Dec.589 (XXVI) of January 2016 on the 1st STC on Legal and Justice Affairs, doc.EX.CL/935 (XXVIII) in which the Assembly adopted the AU Model Law on Medical Products Regulation as an instrument to guide AU Member States in the enactment or review of national medicines laws, and a call to Member States to sign and ratify the said legal instrument, where applicable, as expeditiously as possible to enable its entry into force;

CONVINCED that the efforts to coordinate the regulatory systems strengthening and harmonization initiative under the leadership of African Medicines Agency will provide improved sovereign control and regulation of medical products that will allow African Union Member States to provide for efficient and effective protection of public health against risks associated with use of SF, and will facilitate expeditious approval of products that address the health needs of the African populace, especially for diseases that disproportionately affect Africa.

HAVE AGREED AS FOLLOWS:

PART ONE
THE AFRICAN MEDICINES AGENCY AND ITS OBJECTIVES

ARTICLE 1
ACRONYMS

“AU” refers to the African Union;

“Africa CDC” refers to the Africa Centres for Disease Control and Prevention;

“AMA” refers to the African Medicines Agency;

“AMRC” refers to the African Medicines Regulators Conference;

“AMRH” refers to the African Medicines Regulatory Harmonization Initiative of the African Union;

“API” refers to Active Pharmaceutical Ingredient;

“GMP” refers to Good Manufacturing Practices;

“NEPAD” refers to New Partnership for Africa's Development;

“NMRA” refers to National Medicines Regulatory Authority;

“OAU” refers to Organization of African Unity;

“PMPA” refers to refers to Pharmaceutical Manufacturing Plan for Africa;

“RCOREs” refers to Regional Centres of Regulatory Excellence;

“RECs” refers to Regional Economic Communities recognized by the African Union;

“RHOs” refers to the regional health organizations;

“TC” refers to Technical Committee;

“TWGs” refers to the Technical Working Group comprised of experts constituted under this Treaty;

“WHO” refers to the World Health Organization.

ARTICLE 2 DEFINITIONS

In this Statute, unless the context requires otherwise:

“Agency” means the Agency established under Article 3;

“Assembly” means the Assembly of Heads of State and Government of the African Union;

“Blood Products” means any therapeutic substance prepared from human blood for use in the treatment of diseases or other medical conditions;

“Board” means the Governing Board of the AMA;

“Bureau” means the Bureau of the Conference of the States Parties;

“Commission” means the African Union Commission;

“Complementary Medicines” means any of a range of health therapies that fall beyond the scope of conventional medicine but may be used alongside it in the treatment of diseases and other medical conditions.

“Conference of States Parties” means the Conference of the Parties to this Treaty;

“Constitutive Act” means the Constitutive Act of the African Union;

“Diagnostic” means a medicine or medical device or substance used for the analysis or detection of diseases or other medical conditions.

“Director General” means the Director General of the AMA;

"Food Supplement" means a product intended for ingestion that contains a dietary ingredient intended to add further nutritional value to (supplement) the diet.

"Medical Device" means any instrument, apparatus, implement, machine, appliance, implant, *in vitro* reagent or calibrator, software, material or other similar or related article:

- (a) intended by the manufacturer to be used, alone or in combination, for humans or animals for:
 - (i) diagnosis, prevention, monitoring, treatment or alleviation of disease;
 - (ii) diagnosis, monitoring, treatment, alleviation of or compensation for an injury;
 - (iii) investigation, replacement, modification or support of the anatomy or of a physiological process;
 - (iv) supporting or sustaining life;
 - (v) control of conception;
 - (vi) disinfection of medical devices; or
 - (vii) providing information for medical or diagnostic purposes by means of *in vitro* examination of specimens derived from the human body; and
- (b) which does not achieve its primary intended action in or on the human or animal body by pharmacological, immunological or metabolic means, but which may be assisted in its intended function by such means;

"Medical Products" means medicines, vaccines, blood and blood products, diagnostics and medical devices;

“Medicine” means any substance or mixture of substances used or purporting to be suitable for use or manufactured or sold for use in:-

- (a) the diagnosis, treatment, mitigation, modification or prevention of disease, abnormal physical or mental state or the symptoms thereof in humans; or
- (b) restoring, correcting or modifying any somatic or psychic or organic function in humans, and includes any veterinary medicine;

“Member States” means Member States of the African Union;

“Other Regulated Products” means complementary medicines, traditional medical products, cosmetics, food supplements and related products;

“Secretariat” means the Secretariat of the AMA;

“State Party” means an AU Member State that has ratified or acceded to this Treaty;

“Traditional Medical Product” means an object or substance used in traditional health practice for:

- (a) the diagnosis, treatment or prevention of a physical or mental illness; or
- (b) any curative or therapeutic purpose, including the maintenance or restoration of physical or mental health or well-being in human beings, but does not include a dependence-producing or dangerous substance or drug.

“Treaty” means a treaty to establish the African Medicines Agency.

**ARTICLE 3
ESTABLISHMENT OF THE AMA**

The African Medicines Agency is hereby established as a Specialized Agency of the AU.

**ARTICLE 4
OBJECTIVES OF THE AMA**

The main objective of AMA is to enhance capacity of States Parties and RECs, to regulate medical products in order to improve access to quality, safe and efficacious medical products on the continent.

**ARTICLE 5
GUIDING PRINCIPLES**

The guiding principles of the AMA shall be as follows:

1. **Leadership:** The AMA is an institution that provides strategic direction and promotes good public health practice in States Parties through capacity building, and the promotion of continuous quality improvement in the delivery of medical products regulation;
2. **Credibility:** The AMA's strongest asset is the trust it cultivates with its beneficiaries and stakeholders as a respected, evidence-based institution. It will play an important role in championing effective communication and information-sharing across the continent;
3. **Ownership:** the AMA is an Africa-owned institution. Parties will have primary ownership of AMA to ensure that the financial, human, infrastructural and other resources are adequate for performing its functions;

4. **Transparency and accountability:** The AMA shall operate in accordance with generally accepted international standards of good governance, transparency and accountability:
 - (a) Timely dissemination of information, an open interaction and unimpeded information exchange between the AMA on the one hand, and RECs and Member States on the other;
 - (b) Accountability to States Parties in all its operations;
 - (c) Independent decisions, based on current scientific evidence, professional ethics and integrity. The detailed evidence of its decision-making process and the justification for its decisions shall be fully respected.
5. **Value-addition:** In every strategic aim, objective or activity, the AMA will demonstrate how its initiative adds value to the medical products regulatory activities of States Parties and other partners;
6. **Confidentiality:** The AMA shall adhere to the principles of confidentiality in all its operations;
7. **Commitment to sound quality management:** In all its functions the AMA shall adhere to international standards of quality management and create the conditions for continuous improvement of its regulatory practices and those of NMRAs of Member States of the African Union.

ARTICLE 6
FUNCTIONS

The AMA shall perform the following functions:

- (a) Coordinate and strengthen ongoing initiatives to harmonize medical products regulation and enhance the competence of GMP inspectors to do so;
- (b) Coordinate the collection, management, storage and sharing of information on all medical products including SF medical products, with all its States Parties and globally;
- (c) coordinate joint reviews of applications for the conducting of clinical trials and Provide technical support in quality control of drugs at the request of Member States which do not have the structures to carry out these examination/controls/checks;
- (d) Promote the adoption and harmonization of medical products regulatory policies and standards, as well as scientific guidelines, and coordinate existing regulatory harmonization efforts in the RECs and RHOs;
- (e) Designate, promote, strengthen, coordinate and monitor RCOREs with a view to developing the capacity of medical products regulatory professionals;
- (f) Coordinate and collaborate, where required and on a regular basis, the inspection of drug manufacturing sites, including the regulatory oversight and safety monitoring of medical products, as determined by State Parties and/or the AMA, and make reports available to States Parties;
- (g) Promote cooperation, partnership and recognition of regulatory decisions, in support of regional structures and NMRAs, that takes into account

mobilization of financial and technical resources to ensure sustainability of the AMA;

- (h) Convene, in collaboration with the WHO, the AMRC and other bodies, meetings related to medical products regulation in Africa;
- (i) Provide regulatory guidance, scientific opinions and a common framework for regulatory actions on medical products, as well as priority and emerging issues and pandemics in the event of a public health emergency on the continent with cross border or regional implications where new medical products are to be deployed for investigation and clinical trials;
- (j) Examine, discuss and/or express regulatory guidance on any regulatory matter within its mandate, either on its own initiative or at the request of the African Union, RECs, or States Parties;
- (k) Provide guidance on regulation of traditional medical products;
- (l) Provide advice on the marketing authorization application process for the priority drugs described by the States Parties or on the products proposed by the pharmaceutical laboratories;
- (m) Monitor the medicines market through the collection of samples in every State Party to ensure the quality of selected drugs, have them analysed and provide the results to States Parties and other interested parties, who will thus have reliable information on the quality of the drugs circulating in their countries and, where necessary, will take appropriate measures;
- (n) Develop systems to monitor, evaluate and assess the comprehensiveness of national medical products regulatory systems with the view to recommend measures that will improve efficiency and effectiveness;

- (o) Evaluate and decide on selected medical products, including complex molecules, for treatment of priority diseases/conditions as determined by the African Union, and WHO;
- (p) Provide technical assistance and resources, where possible, on regulatory matters to States Parties that seek assistance and pool expertise and capacities to strengthen networking for optimal use of the limited resources available;
- (q) coordinate access to and network the services available in quality control laboratory services within national and regional regulatory authorities; and
- (r) Promote and advocate for the adoption of the AU Model Law on medical products regulation in States Parties and RECs to facilitate regulatory and legal reforms at continental, regional and national levels.

PART TWO
STATUS OF THE AFRICAN MEDICINES AGENCY AND ITS STAFF

ARTICLE 7
LEGAL PERSONALITY

1. The AMA shall have legal personality that is necessary for the fulfilment of its objectives and the exercise of its functions in accordance with this Treaty;
2. For the smooth fulfilment of its objectives, the AMA shall, in particular, have the legal capacity to:
 - (a) enter into agreements;
 - (b) acquire and dispose of movable and immovable property; and
 - (c) institute and defend legal proceedings.

**ARTICLE 8
PRIVILEGES AND IMMUNITIES**

The General Convention on the Privileges and Immunities of the OAU and the Additional Protocol to the OAU General Convention on Privileges and Immunities, shall apply to AMA, its members, its international personnel, premises, property and assets.

**ARTICLE 9
HEADQUARTERS OF THE AMA**

1. The Headquarters of AMA shall be determined by the Assembly of the Union;
2. The AUC shall enter into a host agreement with the government of the host country in which the AMA Headquarters will be situated with regard to the provision of the premises, facilities, services, privileges and immunities for the purposes of the efficient operation of the AMA.

**PART THREE
ADMINISTRATION AND INSTITUTIONAL FRAMEWORK**

**ARTICLE 10
ORGANS OF THE AMA**

The AMA shall have the following organs:

- (a) The Conference of the States Parties;
- (b) Governing Board;
- (c) The Secretariat; and
- (d) Technical Committees.

**ARTICLE 11
ESTABLISHMENT OF THE CONFERENCE OF THE STATES PARTIES**

The Conference of the States Parties is hereby established as the highest policy-making organ of the AMA. It shall have the power to undertake such functions as are provided for in this Treaty and as may otherwise be necessary to achieve the objectives of this Treaty.

**ARTICLE 12
COMPOSITION OF CONFERENCE OF THE STATES PARTIES**

1. The Conference of the States Parties shall be composed of all Member States of the African Union who ratify or accede to this Treaty;
2. The States Parties shall be represented by Ministers responsible for health or their duly authorised representatives;
3. The Conference of States Parties shall, after due consultation and on the basis of rotation and geographical distribution, elect a Chairperson and other members of the Bureau, namely, three (3) Vice-Chairpersons and a Rapporteur;
4. The Members of the Bureau shall hold office for a period of two (2) years;
5. The Bureau will meet at least once every year;
6. In the absence of the Chairperson or in case of a vacancy, the Vice-Chairpersons or the Rapporteur in order of their election shall act as the Chairperson;

7. The Conference of States Parties shall have the right to invite observers to attend its meetings, and such observers shall not have the right to vote.

**ARTICLE 13
SESSION OF THE CONFERENCE OF THE STATES PARTIES**

1. The Conference of the States Parties shall meet at least once every two years in ordinary session, and in an extraordinary session at the request of the Chairperson, the Bureau, the Governing Board or two-thirds of the State Parties;
2. The quorum of the Conference of the States Parties shall be a simple majority of the States Parties to the AMA;
3. Decisions of the Conference of the States Parties shall be taken by consensus, failing which by a two-thirds majority of the State Parties.

**ARTICLE 14
FUNCTIONS OF THE CONFERENCE OF THE STATES PARTIES**

The Conference of the States Parties shall be responsible for the following functions:

- (a) Set the amount of the annual contribution and special contribution by States Parties, to the budget of the AMA;
- (b) Appoint and dissolve, on good cause, the Governing Board;
- (c) Adopt regulations setting out the powers, duties and conditions of service of the Director General;
- (d) Approve the structure and administrative guidelines of the Secretariat, as well as adopt its governing rules and regulations;

- (e) Provide policy direction to the AMA;
- (f) Recommend the location for the headquarters of the AMA in accordance with the AU criteria adopted by in 2005;
- (g) Approve Regional Centres of Regulatory Excellence (RCORES), on the recommendation of the Governing Board which makes such recommendation after consultation with the Bureau;
- (h) Adopt a scheme to alternate the terms of members of the Board, to ensure that the Board at all times comprises a mix of new and old members;
- (i) Adopt its rules of procedure and for any subsidiary organs;
- (j) Recommend any amendments to this Treaty to the Assembly for consideration.

**ARTICLE 15
ESTABLISHMENT OF THE GOVERNING BOARD**

The Governing Board of the AMA is hereby established by this Treaty. It shall be appointed by and answerable to the Conference of the State Parties.

**ARTICLE 16
COMPOSITION OF THE GOVERNING BOARD**

1. The Board shall consist of Nine (9) members, composed as follows:
 - (a) Five (5) Heads of NMRAs, one (1) drawn from each of the AU-recognized regions;
 - (b) One (1) Representative of RECs responsible for regulatory affairs, to be appointed by the RECs on rotational basis;

- (c) One (1) Representative of Regional Health Organizations responsible for regulatory affairs, on rotational basis appointed by the RHOs;
 - (d) One (1) Representative of National Committees Responsible for Bioethics, on a rotational basis and appointed by the RECs;
 - (e) The Commissioner for Social Affairs, AUC;
2. The Board shall elect its own Chairperson and Vice Chairperson from amongst the Heads of NMRAs;
 3. The Legal Counsel of the AMA or his/her representative shall be an ex-officio member of the Board and shall attend meetings to provide legal advice;
 4. Remuneration for Members of the Board shall be determined by the Conference of the States Parties;
 5. The Director General of the AMA, shall serve as the Secretary of the Board.

**ARTICLE 17
SESSIONS OF THE GOVERNING BOARD**

1. The Board shall meet:
 - (a) in regular session at least once a year;
 - (b) in extraordinary session at the request of the Chairperson of the Board, the Bureau of the Conference of States Parties or a simple majority of the members of the Board;
2. The quorum for meetings of the Board shall be two-thirds of the membership of the Board;

3. The decision of the Board shall be taken by consensus and failing which, by a simple majority vote of the Members present;
4. In the event the Members are not in a position to attend personally, duly accredited representatives shall represent them in accordance with the rules of the governing board;
5. The Board shall consider and recommend its Rules of Procedure and those of the Technical Committees to the Conference of States Parties for adoption;
6. All members of the Board shall be subject to the rules of confidentiality, declaration of interest and conflict of interest;
7. The Board may invite such experts as may be required, to its meetings.

**ARTICLE 18
FUNCTIONS OF THE GOVERNING BOARD**

1. The Board is responsible for providing strategic direction, technical decision-making, guidance and monitoring the performance of the AMA;
2. The functions of the Board shall be to:
 - (a) approve the Strategic Plan, Programme of Work, budgets, activity and reports submitted by the Director General;
 - (b) recommend for endorsement by the Conference of the States Parties, the appointment and dismissal of the Director General of AMA;
 - (c) appoint and dismiss, if necessary, the independent auditor of the AMA;

- (d) recommend regulations setting out conditions of service of the staff of the Secretariat;
- (e) assist the Secretariat with resource mobilization;
- (f) establish technical committees (TCs) to provide technical guidance on the functions of the AMA;
- (g) establish rules governing the issuance of scientific opinions and guidance to States Parties, including expedited approval of products during health outbreaks;
- (h) approve recommendations submitted by the TCs;
- (i) establish such subsidiary or affiliated entities for purposes of carrying out the functions of AMA as it considers necessary;
- (j) carry out any other functions referred to it by the Conference of the States Parties or the Bureau as mandated by the Conference of States Parties.

ARTICLE 19
TERM OF OFFICE OF THE GOVERNING BOARD

1. The term of office of the members of the Board, unless otherwise specified below, shall be a non-renewable period of three (3) years;
2. The term of office of Board members representing the RECs, RHOs shall be a non-renewable period of two (2) years;

3. The Commissioner of Social Affairs (which will become Commissioner for Health, Humanitarian Affairs and Social Development) shall hold a permanent seat;
4. The Board shall elect, by a simple majority and for a three (3) year non-renewable term a Chairperson and Vice Chairperson of the Board from among the heads of NMRA's, taking into account the Union's principle of regional rotation and gender equity.

**ARTICLE 20
ESTABLISHMENT OF TECHNICAL COMMITTEES OF THE AMA**

1. The Board shall establish permanent or ad hoc technical committees to provide technical guidance on specific areas of regulatory expertise;
2. The areas to be considered may include but not be limited to: dossier assessment for advanced therapies, biologicals (including biosimilar and vaccines); medicines for emergencies, orphan medicinal products; clinical trials of medicines and vaccines; manufacturing site inspections of active pharmaceutical ingredients (API) and finished pharmaceutical products, quality control laboratories; bioavailability and bioequivalence studies; pharmacovigilance risk assessment; and African traditional medicines.

**ARTICLE 21
FUNCTIONS OF THE TECHNICAL COMMITTEES**

1. The technical committees shall be responsible for carrying out scientific assessments and conducting scientific reviews of dossiers, including quality aspects, and clinical trial applications; inspection of manufacturing facilities; and providing scientific opinion to facilitate the proper functioning of the AMA;

2. The technical committees shall carry out any other functions as may be assigned to it by the Board.

**ARTICLE 22
COMPOSITION OF THE TECHNICAL COMMITTEES (TCS)**

1. The TCs shall be composed of not more than nine (9) experts representing a wide range of competencies and experiences;
2. Members of the TCs shall be drawn from State Party NMRAs as appointed by the Board and, shall reflect geographic representation;
3. Other technical experts in relevant fields may be drawn from across and outside the continent, when necessary;
4. Each TC shall be headed by a Chair and Vice Chair as specified in its terms of reference adopted by the Board;
5. All members of the TCs shall be subjected to the rules of confidentiality, declaration of interest and conflict of interest.

**ARTICLE 23
THE SECRETARIAT OF THE AMA**

1. The Secretariat of the AMA, located at the headquarters shall be responsible for coordinating the implementation of the decisions of the Conference of the States Parties, the Policy organs of the African Union, and the Board of the AMA;
2. The Secretariat shall:
 - (a) coordinate implementation of activities and ensure effective performance of the AMA in fulfilment of its objectives and functions;

- (b) ensure effective implementation of the decisions of the Board and the Conference of the States Parties;
- (c) coordinate the programmes and work of all technical committees and the Board.
- (d) establish and maintain capacity building and regulatory systems strengthening programmes for the benefit of Member States;
- (e) prepare the strategic plan, work programmes, budget, financial statement and annual report on the activities of the AMA, for consideration and approval by the Board and the Conference of the States Parties;
- (f) perform any other duties as may be assigned by the Board and the Conference of the States Parties and other relevant structures of the African Union.

ARTICLE 24
THE DIRECTOR GENERAL OF THE AMA

1. The Director General shall be the Head of the Secretariat and shall be responsible for the day-to-day management of the AMA;
2. The Director-General shall be appointed by the Conference of the States Parties upon the recommendation of the Governing Board;
3. The Director General, shall serve as the Chief Executive Officer and shall represent the AMA in all matters, and shall report to the Board, the Conference of the States Parties and the African Union, as appropriate;

4. The Director General shall be appointed for a term of four (4) years, renewable once, in accordance with regional rotations;
5. The Director General shall recruit staff of the Secretariat in line with the structure and procedure approved by the Conference of States Parties;
6. The Director General shall be a person of demonstrated competence, leadership ability and integrity, expertise and experience in the subject matter of this Treaty or related issues;
7. The Director General shall be a national of a States Party;
8. The Director General shall be responsible for monitoring the code of conduct of AMA staff and experts;
9. In the discharge of his/her duties the Director General shall not seek or accept instructions from any state, authority or individual external to the AMA.

**ARTICLE 25
OBJECTIONS TO SCIENTIFIC OPINIONS**

1. In the event that a person or entity duly objects to a scientific opinion, advice or decisions issued by AMA, he/she may lodge their objection with the Board;
2. The Board shall set up an independent panel to consider the objection in line with the agreed procedures;
3. The Board shall develop procedures for objection.

**PART FOUR
FINANCIAL PROVISIONS**

**ARTICLE 26
FINANCIAL RESOURCES**

1. The Conference of States Parties shall:
 - (a) set the annual assessed contribution to be paid by the States Parties;
 - (b) adopt the annual the budget of the AMA;
 - (c) determine the appropriate sanctions to be imposed on any Party that defaults in the payment of its contributions to the budget of the AMA in line with the sanctions regime as adopted by the Assembly.
2. The AMA shall devise ways of resource mobilization;
3. The AMA may also receive grants, donations and proceeds for its activities from international organizations, governments, private sector, foundations and other entities in accordance with guidelines set by the Board and approved by the Conference of States Parties, provided there is no conflict of interest;
4. Pending the adoption of the AMA Financial Rules by the Conference of States Parties, it shall abide by the AU Financial Rules and Regulations where appropriate.

**ARTICLE 27
EXPENSES**

1. The Secretariat expenses for administrative, operational and investment purposes shall be in accordance with the approved programme of work,

budget and financial rules and regulations of the AMA as approved by the Governing Board and adopted by the Conference of the States Parties;

2. The finances and accounts of the AMA shall be audited by an independent auditor appointed by the Board.

**PART FIVE
RELATIONS WITH THE AU, MEMBER STATES AND OTHER PARTNER
INSTITUTIONS**

**ARTICLE 28
RELATIONSHIP WITH THE AFRICAN UNION**

1. The AMA shall maintain a close working relationship with the AU;
2. The AMA shall present a written annual report on its activities to the AU Assembly through the relevant STC and Executive Council.

**ARTICLE 29
RELATIONSHIP WITH STATES**

1. The AMA may establish and maintain active cooperation with AU Member States and Non-AU Member States.
2. The States Parties shall appoint focal points to coordinate country level activities of AMA.

**ARTICLE 30
RELATIONSHIP WITH OTHER ORGANIZATIONS AND INSTITUTIONS**

1. The AMA shall establish and maintain a close working relationship and collaboration with the following:

- (a) World Health Organization (WHO);
- (b) Africa Centres for Disease Control and Prevention (Africa CDC);
- (c) Regional Economic Communities (RECs);
- (d) Any other UN agencies, inter-governmental organizations and non-governmental organizations or other institutions, including specialized agencies other than specifically provided for in this Treaty, that AMA considers necessary to assist in achieving its objectives.

**PART SIX
FINAL PROVISIONS**

**ARTICLE 31
WORKING LANGUAGES**

The working languages of the AMA shall be those of the AU, namely Arabic, English, French and Portuguese.

**ARTICLE 32
SETTLEMENT OF DISPUTES**

1. Any dispute that may arise between State Parties with regard to the interpretation, application and implementation of this Statute shall be settled by mutual consent between the States concerned, including through negotiations, mediation, conciliation or other peaceful means;
2. In the event of failure to settle the dispute, the Parties may, by mutual consent, refer the dispute to:

(a) To an Arbitration Panel of three (3) Arbitrators whose appointment shall be as follows:

- i. Each Party to the dispute shall appoint one (1) Arbitrator;
- ii. The third arbitrator, who shall be the Chairperson of the Arbitration Tribunal, shall be chosen by common agreement between the arbitrators appointed by the parties to the dispute; and
- iii. The decision of the Panel of Arbitrators shall be binding.

Or

(b) The African Court of Justice Human and Peoples' Rights.

ARTICLE 33 RESERVATIONS

1. A State Party may, when ratifying or acceding to this statute submit in writing a reservation, with respect to any of the provisions of this treaty;
2. Reservations shall not be incompatible with the objects and purpose of this treaty;
3. Unless otherwise provided, a reservation may be withdrawn at any time;
4. The withdrawal of a reservation must be submitted in writing to the Chairperson of the Commission who shall notify other States Parties of the withdrawal accordingly.

**ARTICLE 34
WITHDRAWAL**

1. At any time after three years from the date of entry into force of this treaty, a State Party may withdraw by giving written notification to the depositary;
2. Withdrawal shall be effective one year after receipt of notification by the depositary, or on such a later date as may be specified in the notification;
3. Withdrawal shall not affect any obligations of the withdrawing State Party prior to the withdrawal.

**ARTICLE 35
DISSOLUTION**

1. The AMA may be dissolved by the agreement of two-thirds of the States Parties to this Treaty at a meeting of the Conference of the States Parties and upon endorsement by the AU Assembly;
2. At least six (6) months' notice shall be given of any meeting of the Conference of the State Parties at which the dissolution of the AMA is to be discussed;
3. Once agreement has been reached on the dissolution of the AMA, the Conference of the States Parties shall establish the modalities for the liquidation of the assets of the AMA.

**ARTICLE 36
AMENDMENT AND REVISION**

1. Any State Party may submit proposals for the amendment or revision of this Treaty. Such proposal shall be adopted at a meeting of a Conference of States Parties;

2. Proposals for amendment or revision shall be submitted to the Chairperson of the Commission who shall transmit the amendment or revision to the Chairperson of the Governing Board within thirty days (30) of receipt thereof;
3. The Conference of States Parties, upon the advice of the Governing Board shall examine these proposals within a period of one year from the date of receipt of such proposals;
4. Amendment or revision shall be adopted by the conference of States Parties by consensus or, failing which, by two thirds majority;
5. The Amendment or revision shall enter into force in accordance with the procedures outlined in Article 38 of this Treaty.

**ARTICLE 37
SIGNATURE, RATIFICATION AND ACCESSION**

1. This Treaty shall be open to Member States of the Union for signature and ratification or accession;
2. The instrument of ratification or accession to the present Treaty shall be deposited with the Chairperson of the Commission who shall notify member states of the union of the deposit of the instrument of ratification or accession.

**ARTICLE 38
ENTRY INTO FORCE**

1. This Treaty shall enter into force thirty days (30) after the deposit of the fifteenth (15) instrument of ratification and accession;
2. The Chairperson of the Commission shall inform all Member States of the Union of the entry into force of the present treaty;

3. For any member state of the Union acceding to the present treaty, the treaty shall come into force in respect of that State on the date of the deposit of its instrument of accession.

**ARTICLE 39
DEPOSITORY**

This Treaty shall be deposited with the Chairperson of the AU Commission, who shall transmit a certified true copy of the Statute to the Government of each signatory State.

**ARTICLE 40
REGISTRATION**

The Chairperson of the Commission upon the entry into force of this Treaty shall register this Treaty with the United Nations Secretary General in conformity with Article 102 of the Charter of the United Nations.

**ARTICLE 41
AUTHENTIC TEXTS**

This Treaty is drawn up in four (4) original texts in the Arabic, English, French and Portuguese languages, all of which are equally authentic.

IN WITNESS WHEREOF, WE the Heads of State and Government or duly authorised representatives of the Member States of the African Union have signed and sealed this Treaty in four original texts in Arabic, English, French, and Portuguese languages, all texts being equally authentic.

**ADOPTED BY THE THIRTY-SECOND ORDINARY SESSION OF
THE ASSEMBLY, HELD IN ADDIS-ABABA, ETHIOPIA**

11TH FEBRUARY 2019

M E M O

TO : DIRECTOR, AUDIT, APPROPRIATION AND
OTHER SELECT COMMITTEES' SERVICES (NA)

FROM : PRINCIPAL CLERK ASSISTANT I

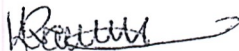
DATE : DECEMBER 5, 2022

SUBJECT : PAPERS LAID

The following Papers were laid on the Table of the House on Thursday, December 1, 2022 (Afternoon Sitting): -

- i.) Memorandum on the Ratification of African Union Treaty for the establishment of the African Medicines Agency (AMA);
- ii.) Annual Report for the Financial Year 2020/2021 from the Parliamentary Service Commission;
- iii.) Reports of the Auditor – General and financial Statements of the following Institutions for the year ended 30th June, 2021 and the certificates therein:
 1. Kitale National Polytechnic;
 2. Bungoma North Technical and Vocational College;
 3. Lugari Diploma Teachers Training College;
 4. Mathioya Technical and Vocational College;
 5. Kenya School of Government;
 6. Institute of human Resource Management;
 7. National Housing Corporation;
 8. Kenya Urban Roads Authority;
 9. Kenya National Highways Authority; and
 10. Lake Basin Development Authority.
- iv.) Reports of the Auditor – General and financial Statements of the following constituencies for the year ended 30th June, 2021 and the certificates therein:
 1. Rangwe;
 2. Suba South;
 3. Likoni;
 4. Kitutu Chache;
 5. South Murirango;
 6. Bobasi;
 7. Mvita;
 8. Malindi;
 9. Magarini;
 10. Awendo;
 11. Nyatibari Chache;
 12. Bonchari;
 13. Taveta; and
 14. Kilifi NORTH

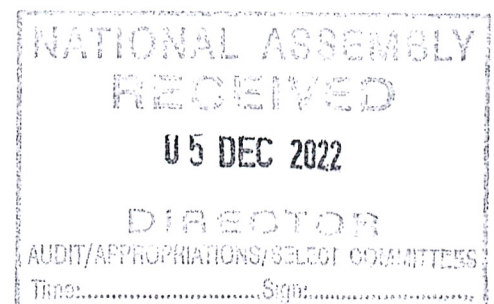
Enclosed herewith, please find the said papers for your necessary action.



RACHEL KAIRU

Copy: Clerk of the National Assembly
Deputy Clerks
Director, Legislative & Procedural Services

(Encls)



SECRET



**REPUBLIC OF KENYA
THE NATIONAL TREASURY AND ECONOMIC PLANNING**

**MEMORANDUM BY THE CABINET SECRETARY FOR THE
NATIONAL TREASURY AND ECONOMIC PLANNING ON THE
CONSIDERATION OF THE RATIFICATION OF THE AFRICAN UNION
TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES
AGENCY BY THE GOVERNMENT OF KENYA**

1. This Memorandum on the consideration of the ratification of the Treaty for the establishment of African Medicines Agency (AMA) by the Government of Kenya; is submitted to the Clerk of the National Assembly by the Cabinet Secretary for the National Treasury and Economic Planning. The Memorandum gives the overview, highlights of the Treaty, and the conclusion.

A. Overview

2. The Treaty seeks to establish the AMA whose main objective will be to enhance capacity of State Parties and Regional Economic Communities (RECs), to regulate medical products in order to improve access to quality, safe and efficacious medical products on the continent. This is based on the premise that access to quality health products and technologies, especially for low and middle-income countries, during the Ebola and COVID-19 pandemics continues to be a challenge due to disruptions in the global supply chain systems. If established, AMA will help African nations to fight pandemics and support national and regional responses by ensuring that only high-quality drugs, vaccines, and other health-related supplies reach African populations.

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B. Highlights of the Treaty

3. The Treaty provides that the functions of the AMA will include but not limited to: providing a platform for coordination and strengthening of on-going regional and continental harmonization of medical products; management of information on all medical products; joint review of clinical trials; adoption and harmonization of medical products regulatory policies and standards; designate promote, strengthen, coordinate and monitor Regional Centers of Regulatory Excellence (RCOREs); coordinate inspection of drugs manufacturing sites; promote partnership of regulatory decisions; convene meetings in collaboration with World Health Organization (WHO); provide regulatory guidance on medical products in Africa; monitor medicines market; promote the adoption of African Union (AU) model law on medical products among others.
4. In relation to the financial implications, the Treaty provides that the Conference of State Parties shall: (a) set the annual assessed contribution to be paid by the States Parties; (b) adopt the annual budget of the AMA; and (c) determine the appropriate sanctions to be imposed on any Party that defaults in the payment of its contributions to the budget of the AMA in line with the sanctions regime as adopted by the Assembly. In addition, the AMA shall devise ways of resource mobilization.
5. Further, the AMA may also receive grants, donations and proceeds for its activities from international organizations, governments, private sector, foundations and other entities in accordance with guidelines set by the Board and approved by the Conference of State Parties, provided there is no conflict of interest. It is further noted that pending the adoption of the AMA Financial Rules by the Conference of States Parties, it shall abide by the AU Financial Rules and Regulations where appropriate.
6. In relation to the expenses, the Treaty provides that Secretariat expenses for administrative, operational and investment purposes shall be in accordance with the approved programme of work, budget and financial rules and

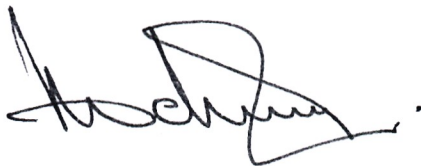
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regulations of the AMA as approved by the Governing Board and adopted by the Conference of the State Parties. In addition, the finances and accounts of the AMA shall be audited by an independent auditor appointed by the Board.

C. CONCLUSION

7. The National Treasury supports the ratification of the Treaty for the establishment of the African Medicines Agencies. The Agency, if established will provide for an African platform for the coordination of regulatory systems regarding medical products. This will ensure that the African population receives quality-assured, safe and efficacious medical products which are fundamental to health and safety for all.



NJUGUNA NDUNG'U, EGH
CABINET SECRETARY

SECRET



MINISTRY OF HEALTH

MEMORANDUM ON RATIFICATION OF THE TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

1.0. OBJECTIVE OF THE MEMORANDUM

- I.1. This Memorandum seeks to support Kenya's ratification of the Treaty for the Establishment of the African Medicines Agency (AMA).

2.0. BACKGROUND

- 2.1. The Treaty seeks to establish the AMA to enhance the capacity of State Parties and Regional Economic Communities (RECs) to regulate medical products to improve access to quality, safe and efficacious medical products on the continent.
- 2.2. AMA is intended to provide a platform for coordination and strengthening ongoing regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources.
- 2.3. The Ministry of Health (MOH), through the Kenya Pharmacy and Poisons Board (KPPB), ensures the **quality, safety and efficacy** of health products and technologies. This has been done through various strategic interventions including building technical capacity, World Health Organisation (WHO) collaborative procedures, reliance mechanisms, harmonisation initiatives and strong collaboration with development partners.
- 2.4. Since 2009, the Africa Union Development Agency (AUDA-NEPAD), working with regional economic communities (RECs)

and collaborating with development partners, has been advancing the African Medicines Regulatory Harmonization (AMRH) Initiative which has now culminated into the Africa Medicines Agency (AMA).

- 2.5. Kenya, led by the Pharmacy and Poisons Board, has been contributing technically to the AMRH initiatives including in the set-up of the EAC-MRH Programme and implementation. These initiatives at the regional and continental levels have to a great extent aided Kenya's realization of its Health Sector development goals and targets while strengthening the national capacity for effective health service delivery.

3.0. CURRENT STATUS OF AMA

- 3.1. As of January 2023, twenty-three (23) member states have fully ratified and deposited their instruments of ratification of the AMA treaty. In **East Africa, Rwanda and Uganda** have **fully ratified** and deposited instruments to the African Union Commission (AUC) while **Tanzania, Burundi and DRC** have **signed** but are yet to ratify and deposit instruments of ratification to AUC.
- 3.2. The AUC is responsible for the administrative and governance setup while AUDA-NEPAD is in charge of the technical set-up of AMA. Currently, **two (2) Kenyans**, officers of the Pharmacy and Poisons Board, are Chairpersons of technical committees under the AUDA-NEPAD AMRH initiative which will be a key precursor of the AMA. i.e. Evaluation of Medicinal Products Technical Committee (EMP-TC) and the Africa Medical Devices Forum (AMDF).
- 3.3. There have been 2 meetings of the Conference of Parties thus far. The Governing Board of the AMA should soon be appointed to enable the appointment of a Director General for the AMA and set up of the Secretariat.
- 3.4. The East African Community is privileged to host AMA headquarters in the Republic of Rwanda.

4.0. OBLIGATIONS IMPOSED ON KENYA BY AMA

- 4.1. Coordinate national and sub-regional medicines regulatory systems;
- 4.2. To conduct regulatory oversight of selected medical products including traditional medicines;
- 4.3. To promote cooperation, harmonisation and the mutual recognition of regulatory decisions;
- 4.4. To strengthen and harmonise efforts of the African Union – recognised RECs, Regional Health Organizations (RHOs) and Member States; and
- 4.5. To complement and enhance collaboration and contribute to improving patients' access to quality, safe and efficacious medical products and health technologies on the continent.

5.0. PROBLEM ANALYSIS

- 5.1. African countries have over-relied on health products manufactured elsewhere despite contributing to 20% of the global burden of disease. Ensuring quality, safe and efficacious health products for the African population remains a core goal of the continent's health products regulators.
- 5.2. The African Pharmaceutical sector is one of the fastest growing in the world and is expected to grow from \$19 billion in 2012 to \$66 billion by 2022. It is estimated that the health and wellness sector in Africa will be worth about \$259 billion by 2030, with the potential to create over 16 million jobs.
- 5.3. Africa has substantively seen a reduction in the number of sub-standard and falsified health products (SFs) circulating in its markets as a result of various joint efforts including in setting up of vibrant Regional Economic Communities (RECs) that work together to review and evaluate applications/dossiers, conduct pharmacovigilance and post-market surveillance across the regions.

- 5.4. Presently, Africa has lagged in regulating complex and specialized molecules, growing its pharmaceutical manufacturing industry and lacks an Active Pharmaceutical Ingredient database that would help enable reliability and trust among its member states.
- 5.5. The COVID-19 pandemic has further triggered the interest of African countries, Kenya included, to develop their manufacturing capacities to remedy the challenges of access to essential health products including vaccines when global supply chains deprioritize Africa's needs.

6.0. JUSTIFICATION FOR RATIFICATION

- 6.1. In the spirit of regionalism and integration, Kenya's role in continental and regional initiatives and being a member of the AU and RECs such as EAC, and IGAD, in signing and ratifying AMA Treaty will demonstrate Kenya's commitment to the Continent's collective action to improved regulation of medicines, medical products and technologies.
- 6.2. **Access to safe, quality and efficacious medical products;**
- (i)** AMA will provide a platform for a multi-faceted approach to combating Substandard and Falsified medical products by strengthening medicine regulatory systems including the capacity for conducting pre-marketing authorizations and routine post-marketing surveillance.
 - (ii)** AMA will complement the National Regulatory Authority's efforts and contribute to capacity building towards improving access to quality-assured medical products within the agenda of Universal Health Coverage and Sustainable Development Goals.
- 6.3. AMA is in line with the Constitution of Kenya, 2010 and will contribute to the achievement of the Kenya Vision 2030 while supporting the achievement of objectives under the Kenya National Health Policy (2018 – 2030), the Kenya Health Sector

Strategic Plan 2018–2023 and the Kenya National Pharmaceutical Policy (KNPP).

6.4. Ease of doing business;

(i) AMA will provide guidance, and streamline and enhance the efforts of REC towards harmonisation of medical products regulation.

(ii) With over 30 pharmaceutical manufacturing plants, Kenya's pharmaceutical industry is the largest in the common market for the Eastern and Southern African regions. AMA is central in ensuring the thriving and development of Africa's Pharma Industry, reducing over-reliance on imported and often expensive medicines and health products.

(iii) AMA will promote local pharmaceutical manufacturing as AMA will reduce duplication of regulatory efforts and ensure efficient use of resources, towards improving access to safe and efficient health products.

6.5. Access to the African Continental Free Trade Area;

(i) In terms of Trade and Economic development, it is anticipated that Kenyan products will have greater access, including reduced time to place products in the markets to a bigger market of all 55 countries in Africa thus benefiting from economies of scale. Additionally, AMA will open up the market for Kenya's local production and manufacturing industry to the USD 1.2 billion markets in Africa, which would go beyond the current USD 160 million at the EAC level.

(ii) AMA will harmonize the regulatory landscape and move Africa towards a truly single integrated block especially when it comes to highly regulated health products. Similarly, AMA will enhance standards, improve the ease of movement of health products that meet accepted international standards, promote local production,

encourage innovation, and ensure efficiency and ease of innovation introduced into the African market.

7.0. CONSTITUTIONAL AND LEGISLATIVE IMPLICATIONS

- 7.1. AMA promotes constitutional values and objectives and does not allude to an amendment to the Constitution.
- 7.2. The Treaty requires AMA to develop, monitor, evaluate and assess the comprehensiveness of National Medical Products Regulatory Systems to recommend measures that will improve efficiency and effectiveness.
- 7.3. The Treaty advocates for the adoption of the African Union Model Law on Regulation of Medical Products to facilitate legal reforms. This may require Kenya to amend its domestic laws to harmonise them with the provisions of the Treaty to facilitate its implementation and to accommodate the work of the AMA.
- 7.4. Kenya may also need to generate guidelines for the periodic reporting obligations generated from joint assessment exercises to establish the capacity of Members States in health products and technologies and technical capacities in line with the proposed logical framework for AMA.

8.0. RESERVATIONS

- 8.1. Article 33 of the AMA Treaty allows ratification with reservations as long as the same is compatible with the objects of the Treaty. There is however no issue in the Treaty that may warrant reservations by Kenya.

9.0. CONCLUSION

- 9.1. All the above-mentioned gains will align with the attainment of Kenya's health priorities for implementation of the Kenya Health Plan (2014-2030), implementation of the Kenya Vision 2030, Sustainable Development Goals (SDGs), Africa Union Agenda 2063 by facilitating access by all citizens to high-quality, safe and efficacious medicines.

9.2. Kenya's growing economy and its comparative advantage across the continent, coupled with its strong pharmaceutical industry, puts the country at a better standpoint to benefit more once AMA comes into force. It is therefore in the national interest that Kenya ratifies the Treaty for the Establishment of AMA.

MEMORANDA ON RATIFICATION OF THE TREATY FOR THE ESTABLISHMENT OF THE AFRICAN MEDICINES AGENCY (AMA)

JUSTIFICATION FOR RATIFICATION

1. Since 2009, the Africa Union Development Agency (AUDA-NEPAD), working with regional economic communities (RECs) and collaborating with development partners, has been advancing the African Medicines Regulatory Harmonization (AMRH) Initiative that has now culminated into the African Medicines Agency (AMA).
2. The AMA is intended to provide a platform for coordination and strengthening on-going regional and continental harmonization initiatives, serving to pool expertise and capacities for optimal use of the limited resources. This memorandum therefore seeks to support Kenya's ratification.
3. AMA will enable Kenya strengthen its Clinical Trials ecosystem including that of COVID-19, strengthen its manufacturing industry, enable it conform to the best practices and standard for health products, strengthen Kenya's capacity to regulate and monitor safety of health products.
4. AMA will provide a platform for a multi-faceted approach for combating Substandard and Falsified medical products by strengthening medicine regulatory systems including the capacity for conducting pre-marketing authorizations and routine post marketing surveillance.
5. The existing national and regional regulatory bodies or harmonization initiatives at RECs level will continue with their mandate but AMA will complement their efforts and contribute to capacity building towards improving access to quality-assured medical products within the agenda of Universal Health Coverage and the Sustainable Development Goals.
6. Kenya supported the AMRH initiatives including in the set-up of the EAC-MRH Programme and implementation. These initiatives at the regional and continental levels have largely aided Kenya's realization of its Health Sector development goals and targets while strengthening the national capacities for effective health service delivery.
7. The establishment of AMA puts Kenya at a better standpoint to benefit more once AMA comes into force. It is therefore in the national interest that Kenya ratifies the Treaty for the Establishment of AMA