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THE KENYA FOOD AND DRUGS AUTHORITY
BILL, 2019

A Bill for

AN ACT of Parliament to establish the Kenya Food and Drugs Authority; to provide for the regulation and management of food, drugs and chemical substances; to provide for the regulation of medical devices and other health technologies; to give effect to the principles and objects of devolved government in food safety regulation and for connected purposes

ENACTED by the Parliament of Kenya, as follows—

PART I—PRELIMINARY

1. (1) This Act may be cited as the Kenya Food and Drugs Authority Act, 2019.

(2) This Act shall come into force on such a date as the Cabinet Secretary may, by notice in the Gazette, appoint, and different dates may be appointed for different provisions.

(3) Subject to subsection (2), no commencement date shall exceed one year from the date of enactment of this Act.

2. (1) In this Act, unless the context otherwise requires—

“advertisement” includes any statement, communication, representation or reference to the public and designed to promote or publicize either directly or indirectly the sale, use or disposal of any health product and technologies including blood and blood products, chemical substances, therapeutic cosmetics, food, herbal medicines and products, medical devices, medicines, scheduled substances, tobacco or tobacco products;

“approved name” in relation to a medicine, means the international non-proprietary name of such medicine or where no such name exists, such other name as the Authority may determine, not being a brand name or trade mark registered in terms of the Trade Marks Act;
“article” includes—

(a) any food, drug, therapeutic cosmetic, herbal medicine, medical device, blood, blood product or scheduled substance and any labelling or advertising materials in respect thereof; or

(b) anything used for the preparation, preservation, packing or storing of any food, drug, herbal medicine, blood, blood product, therapeutic cosmetic, medical device or scheduled substance;

“Authority” means the Kenya Food and Drugs Authority established under section 4 of this Act;

“authorized seller of scheduled substances” means a person designated as such under this Act;

“blood component” means a constituent of blood including erythrocytes, leukocytes, platelets, cryoprecipitate and plasma that can be prepared by various separation methods and under such conditions can be used either directly for therapeutic purposes or for further processing or manufacturing;

“blood product” means any therapeutic substance derived from human blood, including whole blood, blood components and plasma-derived medicinal substances;

“Cabinet Secretary” means the Cabinet Secretary for the time being responsible for matters relating to health;

“chemical substance” means any substance or mixture of substances prepared, sold or represented for use as a germicide; antiseptic; disinfectant; pesticide; insecticide; rodenticide; vermicide; or detergent, or any other substance or mixture of substances which the Authority may, declare to be a chemical substance;

“cigarette” has the meaning assigned to it in the Tobacco Control Act;

“dental practitioner” or “dentist” means a person registered as such under the Medical Practitioners and Dentists Act;

“Director-General” means the Director-General appointed under section 6;
“drug” includes—

(a) any medicine, medicinal preparation, medicinal substance, therapeutic substance or vaccine; or

(b) any substance or mixture of substances including any medicine, medicinal preparation or therapeutic substance prepared, sold or represented for use in—

(i) the diagnosis, treatment, mitigation or prevention of a disease, disorder or abnormal physical state, or the symptoms thereof, in humans or animals; or

(ii) restoring, correcting or modifying functioning if organs in humans or animals;

“enrolled pharmaceutical technologist” has the meaning assigned to it in the Pharmacy and Poisons Act; "falsified medicines" means medicines which do not contain the correct type or concentration of active or other ingredients or falsely labelled medicines;

“food” means any substance, whether processed, semi processed or raw, which is intended for human consumption, and includes drink, chewing gum and any substance which has been used in the manufacture, preparation or treatment of such substances;

“health products and technologies” means blood and blood products, chemical substances, therapeutic cosmetics, food, herbal medicines and products, medical devices including radiation-emitting devices, medicines, scheduled substances, tobacco and tobacco products, and, related products and substances;

“herbal medicine or product” means a plant derived material or preparations with claimed therapeutic or other human or veterinary health benefits, which contain either raw or processed ingredients from one or more plants, or material of inorganic or animal origin;

“insanitary conditions” means such conditions or circumstances as might contaminate food, a drug or a therapeutic cosmetic with dirt or filth or might render the same injurious or dangerous to health;

“interchangeable multi-source medicine” means
medicines that contain the same active substances which are identical in strength or concentration, dosage form and route of administration and meet the same or comparable standards, which comply with the requirements for therapeutic equivalence as prescribed;

"label" includes any legend, work or mark attached to, included in, belonging to or accompanying any food, drug, therapeutic cosmetic, medical device or scheduled substances;

"manufacture" means any process carried out in the course of making a product or medicinal substance and includes packaging, blending, mixing, assembling, distillation, processing, changing of form or application of any chemical or physical process in the preparation of a medicinal substance or product; but does not include dissolving or dispensing the product by diluting or mixing it with some other substances used as vehicle for administration;

"medical device" means any material, instrument, apparatus or contrivance, whether radiation-emitting or not, including components, parts and accessories thereof, manufactured, sold or represented for use in the diagnosis, treatment, monitoring, mitigation or prevention of any disease, disorder or abnormal physical state or disability, or the symptoms thereof, in humans or animals but does not include medicines;

"medicinal substance" means any drug, medicine, product, article, or substance which is claimed to be useful for any of the following purposes—

(a) treating, preventing or alleviating disease or symptoms of disease;
(b) "diagnosing disease or ascertaining the existence, degree or extent of a physiological condition; or
(c) preventing or interfering with the normal operation of a physiological function whether permanently or temporarily and whether by way of terminating, reducing, postponing or increasing or accelerating the operation of the function in human beings or animals;

"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 animals; or

(b) restoring, correcting or modifying any somatic or psychic or organic function in humans or animals;

"medical practitioner" means a person registered as such under the Medical Practitioners and Dentists Act;

"therapeutic cosmetic" includes any substance or mixture of substances manufactured, sold or represented for use in cleansing, improving or altering the complexion, skin, hair, eyes or teeth, and includes deodorants and perfumes;

"tobacco" has the meaning assigned to it in the Tobacco Control Act;

"tobacco product" has the meaning assigned to it in the Tobacco Control Act;

"veterinary surgeon" or "veterinary practitioner" means a person registered as such under the Veterinary Surgeons and Veterinary Para-Professionals Act;

"veterinary medicine" means any curative or preventive substance, formulated medicament, or mixture of substances, whether proprietary or in the form of a preparation effective in animals, which is used, or is manufactured, sold or represented as suitable for use, in—

(a) the diagnosis, treatment, mitigation or prevention of disease or abnormal physical or mental state or the symptoms thereof in an animal; (b) restoring, correcting or modifying any physical, mental or organic function in an animal; or

(c) controlling internal or external pests and parasites, and includes insecticides, vaccines, hormones, alternative medicines, antiseptics, disinfectants, surgical, nutrients and biological products;

"package" includes anything in which any food, drug, therapeutic cosmetic, medical device or scheduled
substances is wholly or partly placed or packed;

"pharmaceutical technologist" has the meaning assigned to it in the Pharmacy and Poisons Act;

"radiopharmaceutical" means a medicinal substance which, when ready for use, contains one or more radionuclides included for a medicinal purpose;

"registered midwife" means a person permitted by law to practice the profession of midwife in Kenya;

"registered pharmacist" means a person registered as such by the body for the time being responsible for registration of pharmacists;

"pharmacy" means either—

(a) the profession of pharmacy as carried out by registered pharmacists; or

(b) the duly licensed premises from which pharmacy services are provided by a registered pharmacist;

"register" means a register established under this Act;

"regulatory officer" means a person appointed by the Authority as such under this Act;

"scheduled substance" means any substance or mixture of substances declared as such in the relevant schedule under this Act;

"serious adverse reaction" means an unforeseen reaction in a blood donor or a recipient associated with collection or transfusion of blood or blood components that has led to death, life-threatening condition, disability, or disease resulting in prolongation of hospitalization;

"substance recommended as a medicine", in relation to the sale of an article consisting of or comprising a substance so recommended, means a substance which is referred to—

(a) on the article or any wrapper or container in which the article is sold, or on any label affixed to, or in any document enclosed in the article, wrapper or container; or

(b) in any placard or other document exhibited at the place where the article is sold; or
(c) in any advertisement published by or on behalf of the manufacturer of the article, or the person carrying on the business in the course of which the article was sold, or, in a case where the article was sold under a proprietary designation, the proprietor of the designation, in terms which are calculated to lead to the use of the substance for the prevention or treatment of any ailment, infirmity or injury affecting human beings or animals, not being terms which give a definite indication that the substance is intended to be used as, or as part of, a food or drink, and not as, or as part of, a medicine.

"substance" includes a preparation or a liquid;

"substandard medicines" means medicines which do not meet defined specifications and includes products that have been contaminated; and

"therapeutic cosmetic" means a therapeutic cosmetic for which a therapeutic effect is claimed;

(2) In this Act, reference to the sale of an article includes reference to the supply of an article as a sample for the purpose of inducing persons to buy by retail the substance of which the article consists or which it comprises.

3. (1) This Act applies to regulation of health products and technologies including—

(a) blood and blood products;
(b) chemical substances;
(c) therapeutic cosmetics;
(d) food;
(e) herbal medicines and products;
(f) medical devices including radiation-emitting devices;
(g) medicines;
(h) scheduled substances; and
(i) tobacco and tobacco products.

(2) Unless otherwise provided in this Act or the Constitution, no other authority or law may regulate the items regulated under this law.
PART II— THE KENYA FOOD AND DRUGS AUTHORITY

4. (1) There is established the Kenya Food and Drugs Authority.

(2) The Authority shall be a body corporate with perpetual succession and a common seal capable, in its corporate name, of—

(a) suing and being sued;
(b) taking, purchasing or otherwise acquiring, holding or disposing of movable or immovable property;
(c) entering into contracts;
(d) borrowing and lending money; and
(e) performing such other things or acts necessary for the proper performance of its functions under this Act which may lawfully be done by a body corporate.

5. The headquarters of the Authority shall be in Nairobi, but the Authority may establish branches anywhere in Kenya.

6. (1) There shall be a Director General of the Authority who shall be appointed by the Public Service Commission through a transparent and competitive process, with the approval of Parliament.

(2) The Director General shall hold office for a term of four years and shall be eligible for reappointment for one further term of four years.

(3) A person shall be qualified for appointment as a Director General if such person—

(a) holds a masters’ degree from a university recognized in Kenya in either pharmacy, food science, medicine or equivalent fields;
(b) has demonstrable experience in the regulation of health products and technologies that affect human health;
(c) has at least ten years’ experience at senior management level in any field relating to the subject matter of this Act;
(d) is a member of a professional body; and
(e) meets the requirements of Chapter Six of the Constitution.

(4) The Director General shall be the chief executive officer of the Authority.

(5) The Director General shall be responsible for the management of the Authority.

(6) The Director General shall be the accounting officer of the Authority.

(7) The Director General shall be the principal representative of the Authority and shall, in that capacity have authority—

(a) to represent the Authority in its relations with other public entities, persons or bodies; and
(b) to sign individually or jointly with other persons, contracts concluded by the Authority, notes and securities issued by the Authority reports, balance sheets, and other financial statements, correspondence and other documents of the Authority.

(8) The Director General may delegate any of his powers provided for in this section to other officers of the Authority.

7. (1) A person shall not qualify for the position of Director General if the person is—

(a) a Member of Parliament;
(b) a Member of a county assembly or county executive committee;
(c) an undischarged bankrupt;
(d) convicted of an offence and sentenced to imprisonment for a term exceeding six months;
(e) a salaried employee of any public entity except on a secondment basis; or
(f) a director, officer, employee, partner in or shareholder of any specified pharmaceutical, food or other institution whose principal business is subject to regulation under this Act.
8. (1) The Authority shall be managed by a Board to be known as the Kenya Food and Drugs Board.

(2) The Board shall comprise—

(a) a non-executive Chairperson to be appointed by the President, through a transparent and competitive process, with the approval of Parliament;

(b) the Principal Secretary in the Ministry for the time being responsible for health or a designated representative;

(c) the Principal Secretary the Ministry for the time being responsible for Finance or a designated representative;

(d) the Director General for Health or a designated representative;

(e) the Director of Veterinary Services or a designated representative;

(f) the Managing Director of Kenya Bureau of Standards or a designated representative;

(g) the Director General of Agriculture, Fisheries and Food Authority or a designated representative;

(h) one person with special knowledge of food sciences nominated by the Council of Governors to represent the interests of counties;

(i) a person representing consumer protection nominated by the body for the time being responsible for consumer protection; and

(j) the Director General who shall be the secretary.

(3) The members of the Board appointed under paragraph (h) and (i) of subsection (2) shall be appointed by the Cabinet Secretary.

(4) A person shall be qualified for appointment to the Board if the person—

(a) is a citizen of Kenya; and

(b) meets the requirements of Chapter 6 of the Constitution.
(5) In addition to the qualifications in subsection (4), the Chairperson to the Board shall possess the following minimum qualifications—

(a) a masters degree in science from a recognized university; and

(b) at least 15 years experience in any field relating to the subject matter of this Act.

(6) A person shall not be qualified for appointment as member of the Board if such person—

(a) is a member of Parliament;

(b) is a member of a county assembly or executive;

(c) is a member of a governing body of a political party;

(d) is an undischarged bankrupt; or

(e) is convicted of an offence and sentenced to imprisonment for a term exceeding six months.

(7) In appointing members of the Board, regard shall be had of the need for regional balance and the realisation of the principle that at not more than two thirds of the members shall be of the same gender.

(8) The members shall, at their first meeting, elect a Vice-chairperson from amongst the members appointed under sub-sections (2)(b) to (i).

(9) The Chairperson and the Vice-chairperson of the Board shall not be of the same gender.

(10) The members of the Board shall hold office for a term of three years and shall be eligible for reappointment for one further term of three years.

(11) The members of the Board shall be appointed in accordance with the Third Schedule to this Act.

9. The Chairperson and all the members of the Board shall subscribe before the Chief Justice the oath or affirmation set out in the Second Schedule to this Act.

10. (1) The office of the Chairperson or of a member of the Board shall become vacant if the holder—

(a) dies;
(b) resigns from office by writing under his hand addressed to the President;

c) is removed from office in accordance with the provisions of section 11;

d) is convicted of an offence and sentenced to imprisonment for a term exceeding six months without the option of a fine;

(e) is unable to discharge the functions of his office by reason of physical or mental infirmity;

(f) is absent without permission of the Chairperson from three consecutive meetings of the Board without good cause; or

(g) is declared bankrupt.

(2) A Board member may resign from the position of Vice chairperson without losing his or her position as Board member.

11. (1) The Chairperson or a member of the Board may be removed from office for—

(a) gross violation of the Constitution or any other law;

(b) gross misconduct, whether in the performance of the member's functions or otherwise;

(c) physical or mental incapacity to perform the functions of office; or

(d) incompetence or neglect of duty.

(2) The Cabinet Secretary may, upon the recommendation of the Board, revoke the appointment of a member of the Board on any of the grounds specified under sub-section (1).

12. The primary object of the Authority is to provide for the regulation, investigation, inspection and approval of food, health products and technologies, and related matters in the public interest, and for that purpose the Authority shall—

(a) ensure adequate and effective standards and guidelines for regulation of health products and technologies;
(b) ensure that compliance with existing legislation is being promoted and controlled through a process of active inspection and investigation;

(c) ensure the efficient, effective and ethical evaluation and registration of health products and technologies that meet defined standards of quality, safety and efficacy;

(d) ensure that the process of evaluating and registering health products and technologies is, subject to this Act, transparent, fair, objective and concluded timeously;

(e) ensure the periodic re-assessment and monitoring of health products and technologies;

(f) ensure that evidence of existing and new adverse events, interactions, information about health products and technologies is being monitored globally, analysed and acted upon;

(g) ensure that clinical trial protocols where required for registration of health products and technologies are being assessed according to prescribed ethical and professional criteria and defined standards;

(h) monitor compliance with this Act through its agencies and any other agencies of State authorised under this Act;

(i) advise the cabinet secretary and county governments on measures for the protection of the health of consumers;

(j) advise the cabinet secretary and county governments on the implementation of this Act;

(k) foster co-operation between the Authority and other institutions or organizations and other stakeholders;

(l) approve and register health products and technologies regulated under this Act, manufactured within or imported into, and intended for use in Kenya;

(m) examine, grant, issue, suspend, cancel and revoke licences or permits issued under this Act;
(n) appoint inspectors and order inspection of any premises;
(o) promote rational use of drugs, medical devices and herbal drug;
(p) provide the public with unbiased information on products regulated under this Act;
(q) prescribe standards of quality in respect of products regulated under this Act, manufactured or intended to be manufactured or imported into or exported from the Kenya;
(r) maintain registers prescribed under this Act;
(s) attend to and, where possible, take legal measures on complaints made by consumers against manufacturers of products regulated under this Act;
(t) be responsible for its human resource management and development; and
(u) perform any other functions assigned to it under this Act.

13. The Authority shall have powers necessary or expedient for the proper performance of its functions under this Act including—

(a) to enter into association with such other bodies or organizations within or outside Kenya as it may consider desirable or appropriate and in furtherance of the purpose for which the Authority is established;
(b) control, supervise and administer the assets of the Authority in such manner and for such purposes as best promote the purpose for which the Authority is established;
(c) receive any grants, gifts, donations or endowments and make legitimate disbursements there from;
(d) open banking accounts for the funds of the Authority;
(e) establish such committees as it may consider necessary for the performance of its functions and the exercise of its powers under this Act; and

Powers of the Authority.
(f) co-opt in such committees persons whose knowledge and experience is necessary to enable to the committee effectively to discharge its functions.

14. The conduct and regulation of the business and affairs of the Board of the Authority is provided in the First Schedule to this Act.

15. The Board may, by resolution either generally or in any particular case, delegate to any committee of the Authority or to any member, officer, employee or agent of the Authority, the exercise of any of the functions or duties of the Authority under this Act.

16. The cabinet secretary shall, on the advice of the Salaries and Remuneration Commission, determine the allowances of the members of the Board of the Authority.

17. (1) The Authority may appoint such officers or staff including regulatory officers, as are necessary for the proper discharge of the functions of the Authority under this Act, upon such terms and conditions of service as the Authority may determine.

(2) The principles of ethnic, regional and gender balance shall guide all staff appointments.

(3) The Cabinet Secretary may, upon request by the Authority, second to the Authority such number of public officers as may be necessary for the purposes of the Authority.

(4) A public officer seconded to the Authority shall, during the period of secondment, be deemed to be an officer of the Authority and shall be subject only to the direction and control of the Authority.

18. (1) The common seal of the Authority shall be kept in such custody as the Authority may direct and shall not be used except on the order of the Board.

(2) The common seal of the Authority when affixed to a document and authenticated shall be judicially and officially noticed and unless and until the contrary is proved, any necessary order or authorization of the Board under this section shall be presumed to have been given.

19. Nothing done by a member of the Board or any officer, employee or agent of the Authority shall, if the matter or thing is done in good faith for executing the
functions, powers and duties of the Authority render the member, officer, employee or agent personally liable to any action, claim or demand whatsoever.

20. (1.) The Director General shall be the Registrar of the Authority.

(2) The Registrar shall perform such duties and exercise such powers, in addition to those required under the provisions of this Act to be performed and exercised, as the Board may from time to time direct.

21. (1) The Board may establish scientific advisory committees, as may be necessary for the performance of the Board’s functions and the exercise of the Board’s powers under this Act.

(2) The scientific advisory committees established under subsection (1) shall be in addition to those established in the Fourth Schedule to this Act.

(3) The primary role of the advisory committees shall be to provide the Board with expert, independent advice on complex scientific issues presented to the Authority.

(4) The Board may co-opt into the membership of committees established under subsections (1) and (2) above, other persons whose knowledge and skills are found necessary for the functions of the Authority.

(5) A committee may be so established for any purpose, or combination of purposes, connected with the execution of this Act or the exercise of any power conferred to it, either generally or in relation to any particular class of substances or articles to which any provision of this Act is applicable.

(6) Without prejudice to the generality of subsection (3) of this section, in relation to any such class of substances or articles, a committee may be established under this section for either or both of the following purposes—

(a) giving advice with respect to safety, quality or efficacy; or

(b) promoting the collection and investigation of information relating to adverse reactions, for the purpose of enabling such advice to be given.
(7) The Authority shall provide adequate staff to enable the advisory committees to perform their functions effectively.

(8) The chairperson of an advisory committee shall convene a meeting of the Committee at least once every two months and shall convene an additional meeting if requested by at least four members in writing.

(9) An advisory committee shall submit, at least once every six months, a report to the Cabinet Secretary, with respect to its activities and the Cabinet Secretary shall lay a copy of each report before the Parliament.

(10) The quorum of an advisory committee shall be five members with one member being appointed as chairperson.

PART III—FOOD

22. (1) The Authority shall regulate and monitor the manufacture, processing, distribution, warehousing, wholesale and importation of food in Kenya.

(2) Without prejudice to the provisions of sub-section (1), the Authority shall—

(a) set the standards and guidelines in relation to the regulation of food products and food additives;

(b) provide scientific advice and technical support to the National Government and the county governments in matters of framing the policy and rules in areas which have a direct or indirect bearing on food safety and nutrition;

(c) search, collect, collate, analyse and summarise relevant scientific and technical data particularly relating to—

(i) food consumption and the exposure of individuals to risks related to the consumption of food;

(ii) incidence and prevalence of biological risk;

(iii) contaminants in food;

(iv) residues of various contaminants;

(v) identification of emerging risks; and

(vi) introduction of rapid alert system;
(d) promote, co-ordinate and issue guidelines for the development of risk assessment methodologies;

(e) establish systems and mechanisms for the reporting of adverse reactions to food;

(f) establish mechanisms for the disposal and cancellation of licences;

(g) provide scientific and technical advice and assistance to the National Government and the county governments in implementation of crisis management procedures with regard to food safety;

(h) draw up a general plan for crisis management and work in close co-operation with the county based crisis units;

(i) establish a system of network of organisation with the aim to facilitate a scientific co-operation framework by the co-ordination of activities, the exchange of information, the development and implementation of joint projects, the exchange of expertise and best practices in the fields within the Authority's responsibility;

(j) provide scientific and technical assistance to the national government for improving co-operation with international organisations;

(k) take all such steps to ensure that the public, consumers, interested parties receive rapid, reliable, objective and comprehensive information on food safety through appropriate methods and means;

(l) provide, whether training programmes in food safety and standards for persons who are or intend to become involved in food businesses, whether as food business operators or employees or otherwise; and

(m) promote general awareness as to food safety and food standards.

23. (1) A person shall not sell food that—

(a) Contains any poisonous or harmful substance;

(b) is unwholesome or unfit for human consumption;

Prohibition against sale of unwholesome, poisonous or adulterated food.
(c) prohibited under this Act or any other law;
(d) contains any filthy, putrid, rotten, decomposed or diseased substance or foreign matter; or
(e) is adulterated.

(2) A person who contravenes sub-section (1) commits an offence.

(3) Subsections (1) and (2) shall apply in relation to—

(a) any food intended for human consumption which is offered as a prize or reward or donation in connection with any entertainment to which the public are admitted, whether or not on payment of money, as if such food were or had been, exposed for sale by each person in the organization of the entertainment;

(b) any food intended for human consumption which is offered as a prize or reward, donation or given away for the purposes of advertisement, or in furtherance of any trade or business, as if the food were, or had been, exposed for sale by the person offering or giving it away; and

(c) any food intended for human consumption which is exposed or deposited in any premises for the purpose of being so offered or given away as if the food were, or had been, exported for sale by the occupier of the Premises.

24. (1) A person who has reasons to believe that a food which the person has processed, manufactured or distributed is not in compliance with this Act, any other law, the person shall immediately initiate procedures to withdraw the food in question from the market and consumers indicating reasons for its withdrawal and inform the Authority and the relevant food safety authority at the county level.

(2) A person shall immediately inform the Authority and the relevant food safety authority at the county level co-operate with the authorities, if the person considers or has reasons to believe that a food which he has placed on the market may be unsafe for the consumers.

(3) The person under subsections (1) and (2) shall inform the authorities of the action taken to prevent risks to the consumer and shall not prevent or discourage any Duty to notify authorities.
person from co-operating, in accordance with this Act, with the competent authorities, where this may prevent, reduce or eliminate a risk arising from a food.

(4) Every person shall follow such conditions and guidelines relating to food recall procedures as the Authority may specify by regulations.

25. A person who labels, packages, treats, processes, sells or advertises any food in contravention of this Act or any other law, or in a manner that is false, misleading or deceptive as regards its character, nature, value, substance, quality, composition, merit or safety, commits an offence.

26. A person who labels, packages, sells or advertises any food in contravention of a standard prescribed under this Act commits an offence.

27. A person who any food which is not of the nature, substance, or quality demanded by the purchaser commits an offence.

28. A person who sells, prepares, packages, conveys, stores or displays for sale any food under insanitary conditions commits an offence.

29. A person shall not import into Kenya—

(a) any unsafe or misbranded or sub-standard food;

(b) any article of food the import of which a licence is required under this Act or any other law except in accordance with the conditions of the licence;

(c) which contains or has been treated with a prohibited substance;

(d) which contains a particular substance in a greater measure than that permitted under this Act or any other law or has been treated with a substance containing a particular substance in a greater measure than that permitted by regulation;

(e) which has been treated in such manner that its damaged or unsound condition or inferior quality is concealed whether entirely or partly; and

(f) any article of food in contravention of this Act or any other law.
30. (1) A regulatory officer may, examine any food intended for human consumption which has been distributed, sold, or is offered or exposed for sale or is in the possession of, or has been deposited with or consigned to, any person for the purpose of distribution or sale or manufacture for sale, if it appears to the regulatory officer to be unfit for human consumption, may seize it and remove it in order to have it dealt with in a manner provided for in this Act.

(1) A regulatory officer who seizes any food under subsection (1) shall inform the person in whose possession the food was found of the seizure of the food and the intention to have the food tested.

(2) If it appears to the regulatory officer that any perishable food examined by the regulatory officer is unfit for human consumption, the regulatory officer shall confiscate the food and in the prescribed manner, transmit it to the laboratory for testing.

(3) When it appears to the regulatory officers that any non-examined perishable food is unfit for human consumption the regulatory officer shall take that food to the court for further action, and if the court is satisfied that, that food is unfit for human consumption, the court shall order for its destruction or disposal in a prescribed manner, and if the court is not satisfied that the food is unfit for human consumption, it may order for further actions to be taken.

(4) If the court orders for the destruction or disposal of any food which has been declared unfit for human consumption, that destruction or disposal shall be done at the owners cost.

(5) authorized regulatory officer may seize, and cause to be disposed of in the prescribed manner, the carcass or any part of a carcass found in a butchery facility or cold stores for the purpose of sale for human consumption which on examination in the prescribed manner is found to be diseased or unfit for human consumption, and no compensation shall be payable to the owner of that carcass or any part of it.

31. Any food which either contains a scheduled substance or which claims to have a therapeutic effect or value shall be treated as a medicine under this Act.
32. (1) There is established under this Act, a Therapeutic Foods Register.

(2) The Registrar shall have custody of the Register.

(3) The register kept under subsection (1) shall be available for inspection by the public.

(4) The Authority shall prescribe the format and the content of Therapeutic Foods Register.

PART IV—MEDICINES

33. (1) A person shall not sell any medicine that—

(a) is not registered by the Authority;

(b) is adulterated;

(c) is substandard; or

(d) fails to comply with any specifications made under this Act or any other written law.

(2) A person who contravenes the provisions of subsection (1) commits an offence and upon conviction, is liable—

(a) in the case of a first offence, to a fine not exceeding one million shillings or to imprisonment for a term not exceeding three months, or to both; or

(b) in the case of a subsequent offence, to a fine not exceeding two million shillings or to imprisonment for a term not exceeding five years, or to both.

(3) Sub-section (1) shall not apply to the sale of a medicine compounded by a pharmacist or pharmaceutical technologist—

(a) in a quantity not greater than that prescribed under this Act for sale in the retail trade, subject to prescribed conditions; or

(b) in a quantity for a particular person or animal as prescribed by an approved medical practitioner as the case may be, if that pharmaceutical product does not contain any component the sale of which is prohibited by this Act, or any component in respect of which an application has been rejected,
and if that pharmaceutical product has not been advertised.

34. (1) A person shall not—
(a) falsify medicines;
(b) label, package, treat, process, sell or advertise any medicine in contravention of any regulations made under this Act; or
(c) make statements regarding the character, constitution, value, potency, quality, composition, merit or safety of a medicine in a manner that is false, misleading or deceptive.

(2) A person who contravenes subsection (1) above commits an offence and upon conviction, is liable—
(a) in the case of a first offence, to a fine not exceeding one million shillings or to imprisonment for a term not exceeding three years or to both; or
(b) in the case of a subsequent offence, to a fine not exceeding two million shillings or to imprisonment for a term not exceeding five years, or to both.

35. (1) If a standard has been prescribed for a medicine, a person who manufactures, labels, packages, sells or advertises any substance in such a manner that it is likely to be mistaken for that medicine having met the prescribed standard commits an offence unless the substance is the medicine in question and complies with the prescribed standard.

(2) If a standard has not been prescribed for a medicine but a standard for the medicine is contained in any of the publications specified in the Fifth Schedule, any person who manufactures, labels, packages, sells or advertises any other substance or article in such a manner that it is likely to be mistaken for the drug having met any of the standards contained in any of the publications specified in the Fifth Schedule commits an offence.

(3) A person who manufactures, labels, packages, sells or advertises any medicine for which no standard has been prescribed, or for which no standard is contained in any of
the publications specified in the Fifth Schedule, commits an offence unless the medicine—

(a) is in accordance with the professed standard under which it is labelled, packaged, sold or advertised; and

(b) does not resemble, in a manner likely to deceive, any drug for which a standard has been prescribed or which is contained in any of the publications specified in the Fifth Schedule.

(4) A person convicted of an offence under subsections (1), (2) and (3) is liable in the case of a first offence, to a fine not exceeding one hundred thousand shillings or to imprisonment for a term not exceeding three months, or to both such fine and imprisonment and in the case of a subsequent offence, to a fine not exceeding two hundred thousand shillings or to imprisonment for a term not exceeding five years, or to both.

36. A person who sells any medicine which is not of the nature, substance, or of the quality, of the article demanded by the purchaser commits an offence and upon conviction, is liable—

(a) in the case of a first offence, to a fine not exceeding one one million shillings or to imprisonment for a term not exceeding three years, or to both such fine and imprisonment; or

(b) in the case of a subsequent offence, to a fine not exceeding two million shillings or to imprisonment for a term not exceeding five years, or to both such fine and imprisonment.

37. A person who manufactures, sells, prepares, preserves, packages, stores or conveys for sale any medicine under conditions not meeting prescribed standards commits an offence.

38. In considering an application for a product licence the Authority shall in particular take into consideration—

(a) the safety of the medicinal products to which the application relates;

(b) the efficacy of the medicinal products to the purpose for which they are proposed to be administered;
(c) the quality of the medicinal products of each such description, according to the specification and the method or proposed method of manufacture of the products, and the provisions proposed for securing that the products as sold or supplied will be of that quality; and

(d) any other factor that the Authority considers necessary.

39. (1) There is established under this Act, a Medicines Register.

(2) The Authority shall prescribe the format and the content of the Medicines Register.

40. (1) Every application for the registration of a medicine or medical device shall be submitted to the Registrar in the prescribed form and shall be accompanied by the prescribed particulars and samples of the relevant medicine and by the prescribed registration fee.

(2) The Registrar shall ensure that such an application in respect of medicine which appears on the latest Essential Medicines List or Essential Veterinary Medicines List or a medicine which does not appear thereon but which, in the opinion of the relevant cabinet secretary, is essential for national human or veterinary health is subject to such procedures as may be prescribed in order to expedite the registration.

(3) After consideration of an application and after any investigation or enquiry which it may consider necessary, the Authority shall approve of the registration if the authority is satisfied that—

(a) the medicine in question is suitable for the purpose for which it is intended;

(b) the medicine in question complies with the prescribed requirements; and

(c) that registration of that medicine is in the public interest.

(4) If the Authority does not approve the registration of a medicine, it shall cause the applicant to be notified:

(a) in writing of the reasons why it is not so satisfied; and
(b) that the applicant may within a period of one month after the date of the notification furnish the registrar with the applicant's reasons for not being so satisfied.

(5) If no comments under subsection (4) above are submitted by the applicant within the prescribed period, or if after consideration of any comments so submitted the Authority is still not satisfied as aforesaid, the authority shall reject the application.

(6) When the Authority has approved of the registration of any medicine, the registrar shall register that medicine and shall enter in the register such particulars in regard to the medicine as are required by this Act to be so entered and shall issue to the applicant a certificate of registration in the prescribed form in respect of that medicine.

(7) Every medicine shall be registered under such name as the Authority may approve.

(8) The Registrar shall allocate to every medicine registered under this Act a registration number which shall be recorded in the register opposite the name of such medicine and which shall be stated in the certificate of registration issued in respect of such medicine.

(9) Any registration under this section, including the registration of medicines already registered, shall be valid for a period of five years and may be made subject to such conditions as may with regard to the succeeding provisions of this section be determined by the Authority.

(10) No condition shall be imposed under sub-section (9) whereby the sale of the medicine in question by any person other than a pharmacist is prohibited or until after the applicant has been notified in writing by the Registrar that the imposition of such condition is contemplated and invited to submit written representations to the Authority in regard to the matter.

(11) If no representations under sub-section (10) above are lodged with the Registrar by the applicant concerned within a period of one month after the receipt by of the notification referred to in sub-section (10), or if after consideration of any such representations the Authority is still of the opinion that the condition in question should be
imposed, the Authority shall direct the registrar to register the medicine concerned subject to the said condition.

(12) Notice of the rejection of an application under this section in respect of a medicine referred to in subsection (4) shall be given in the gazette by the registrar-

(a) if no appeal is lodged against the rejection within the sixty days, as soon as possible after the expiration of that period; or

(b) if any appeal so lodged is dismissed, as soon as possible after the decision dismissing the appeal has been given.

(13) The Registrar shall as soon as possible after the date of expiry of the appropriate period referred to in sixty days publish in the Gazette the prescribed particulars in respect of all applications for registration received by him or her prior to such date.

(14) For the purposes of this section—

(a) Essential Medicines List means the list of essential medicines included in the latest edition of the official publication relating to guidelines for standard treatment which is compiled by the State Department responsible for Health; and

(b) Essential Veterinary Medicines List means the list of essential medicines included in the latest edition of the official publication relating to guidelines for standard treatment which is compiled by the State Department responsible for Veterinary Medicines.

41. (1) The Registrar, on application by the holder of a certificate of registration issued in respect of a medicine and with the approval of the Authority, may amend the entry in a register with respect to that medicine.

(2) An application for the amendment of an entry in a register shall be made to the registrar in the prescribed form and shall be accompanied by the prescribed application fee.

(3) If the Authority grants its approval in respect of an application submitted to it in terms of subsection (2), the registrar shall—
(a) make the required amendments in the relevant register; and

(b) if the name of the applicant changes, issue a new certificate of registration on the prescribed form to the applicant in respect of the medicine after receiving the existing certificate of registration in respect of that medicine for cancellation.

42. (1) The holder of a certificate of registration may transfer, with the approval of the authority, the certificate of registration to another person, who is duly licensed to practice the profession of pharmacy and holds a valid practising certificate to apply for the registration of a medicine.

(2) An application for approval of the transfer of a certificate of registration shall be made to the registrar in the prescribed form and shall be accompanied by the certificate of registration in question and the prescribed application fee.

(3) If the Authority allows the application submitted to it in terms of subsection (2) above, the registrar shall—

(a) make the necessary entries in the register relating to the person to whom the certificate of registration is transferred;

(b) cancel the existing certificate of registration; and

(c) issue a new certificate of registration on the prescribed form to the person making the application in respect of the relevant medicine.

43. (1) The Authority shall cancel the registration of a medicine or medical device if—

(a) a licencee has failed to comply with a condition subject to which a particular medicine or medical device has been registered;

(b) a particular medicine or medical device does not comply with a prescribed requirement; or

(c) it is not in the public interest to make a particular medicine or medical device available to the public.

(2) Before cancellation of the registration of any
(3) The notice referred to in subsection(2) above shall—

(a) specify the grounds on which the decision of the Authority is based; and

(b) indicate that the person to whom the notice is directed may within one month after the date of that notice submit to the Registrar any objection, which he or she may wish to make in connection with the matter.

(4) The Authority shall direct the Registrar to cancel the registration of that medicine or medical device, if—

(a) no objection as contemplated in subsection (2)(ii) are received; or

(b) after consideration of any comments received, the Authority is of the opinion that the registration of the medicine or medical device in question should be cancelled.

(5) If the holder of the certificate of registration issued in respect of a medicine or medical device fails to pay the prescribed fee in respect of the retention of the registration of that medicine or medical device before or on the prescribed date or such later date as the registrar, with the approval of the Authority, may determine on application by that person, the registrar shall cancel the registration of that medicine or medical device.

44. (1) The registrar shall give notice in the gazette of the registration, or cancellation of registration, of a medicine or medical device in terms of this Act.

(2) The Registrar shall in such notice specify in the case of a registration of a medicine—

(a) the name under which that medicine is registered;

(b) the active components of that medicine;

(c) the name of the applicant;

(d) the name of the manufacturer;
(e) the registration number allocated to that medicine and
(f) the conditions, if any, subject to which that medicine is registered;

(3) In the case of a cancellation of registration of a medicine medical device the Registrar shall in such notice specify—
(a) the name under which that medical device was registered;
(b) the name of the holder of the certificate of registration issued in respect of that medical device; and
(c) the number which was allocated to that medical device in terms of this Act.

45. The Cabinet Secretary in consultation with the authority, may prescribe conditions for the supply of more affordable medicines in certain circumstances so as to protect the health of the public, and in particular may—
(a) prescribe the conditions under which any medicine which is identical in composition, meets the same quality standard and is intended to have the same proprietary name as that of another medicine already registered in the Republic, but which is imported by a person other than the person who is the holder of the registration certificate of the medicine already registered and which originates from any site of manufacture of the original manufacturer as approved by the Authority in the prescribed manner, may be imported; or
(b) prescribe the registration procedure for, as well as the use of, the medicine referred to in paragraph (a).

46. (1) A pharmacist shall dispense an interchangeable multi-source medicine instead of the medicine prescribed by a medical or dental practitioner, nurse or other person registered under the relevant statutes regulating health professionals.

(2) A medical or dental practitioner, nurse or other person registered under the relevant statutes regulating
health professionals may prohibit a pharmacist is from interchanging a medicine contemplated in subsection (1) above and that fact shall be noted on the prescription.

(3) When an interchangeable multi-source medicine is dispensed by a pharmacist he or she shall note the brand name or where no such brand name exists, the name of the manufacturer of that interchangeable multi-source medicine in the prescription book.

(4) A pharmacist shall not substitute for a prescribed medicine an interchangeable multi-source medicine—

(a) if the person prescribing the medicine has written in his or her own hand on the prescription the words 'no substitution' next to the item prescribed;

(b) if the retail price of the interchangeable multi-source medicine is higher than that of the prescribed medicine; or

(c) where the product has been declared not substitutable by the Authority.

47. (1) A person who, on a commercial scale, manufactures, prepares, supplies, sells, distributes, exports or imports a herbal medicine which is represented to the public to have therapeutic effect when it is consumed, applied or inhaled must be licensed.

(2) Regulations may prescribe particulars to be provided for in the registration of items under sub-section (1).

(3) A person who contravenes sub-section (1) or any other law commits an offence.

PART V—SCHEDULED SUBSTANCES

48. (1) The Authority shall prepare and submit to the Cabinet Secretary lists of the substances which are to be treated as Scheduled Substances for the purposes of this Act.

(2) The lists to be prepared under this Section shall include—

(a) substances which, subject to this Act, are not to be sold except by authorized sellers of Scheduled
Substances and by licenced wholesale dealers and dealers in mining, agricultural or horticultural accessories;

(b) substances which, subject to this Act, are not to be sold except by persons specially licensed to do so; and

(c) any other substance declared to be a Scheduled Substance by the authority.

(3) The cabinet secretary may, by order, confirm the list with or without modification, and may, after consultation with or on the recommendation of the Authority, by order amend or vary the list.

(4) The authority shall review the lists under subsection (2) at least once every year.

49. (1) The following persons may be in possession of Scheduled Substances, but to the extent only and subject to the Limitations prescribed by this sub-section—

(a) a wholesale dealer licensed under this Act, for the purposes of the licence and on the premises so licensed;

(b) an authorized seller of Schedules Substances, on premises registered by the Authority;

(c) a person licensed by the Authority to sell Schedules Substances for mining, agricultural or horticultural purposes, for the purposes of the licence and on premises so licensed;

(d) a person, institution or department, to which a Scheduled Substance has been lawfully sold under this Act, for the purpose for which the sale was made;

(e) a person for whom the Scheduled Substance has been lawfully supplied or dispensed by a qualified medical practitioner, dentist or veterinary surgeon, or by a hospital, dispensary or similar institution;

subject to any conditions which may be prescribed, a representative of the person engaged in the business of selling and supplying pharmaceutical goods, for the
purpose of giving free samples of those goods, in the course of the business, to persons who may lawfully be in possession of Scheduled Substances;

(f) the personal representative of a deceased person, or the liquidator, receiver or other person appointed to deal with the property of a bankrupt or of a company which is being wound up compulsorily, or the manager of the estate of a person of unsound mind, in respect of poisons in the possession of the deceased person, bankrupt person, company or person of unsound mind at the time of death or bankruptcy or the beginning of the winding up of the order appointing the manager, for the purpose of disposing up of the Scheduled Substances, with the written permission of the Authority and in accordance with its directions, to a wholesale dealer in Scheduled Substances licensed under this Act or to an authorized seller of Scheduled Substances.

(2) A person who is in possession of a Scheduled Substance otherwise than in accordance with the provisions of this Section shall commit an offence and upon conviction, is liable to a fine not exceeding one hundred thousand shillings or to imprisonment for a term not exceeding three years or to both.

50. (1) If the Authority is satisfied that it is in the public interest that a licence to deal as a wholesale dealer in Scheduled Substances should be issued or renewed it may, on application being made to the Authority in writing on such form as may be prescribed, and on payment of the prescribed fee, issue to the applicant a licence in the form prescribed, or, as the case may be, renew the licence.

(2) The Authority may refuse to issue or renew, or may revoke, a licence under this Section, for any good and sufficient reason relating either to the applicant or licensee, or to the premises in which the business is, or is proposed to be, carried on, and in case of refusal or revocation an appeal shall lie to the Cabinet Secretary.

(3) A separate licence under this Section shall be required in respect of each set of premises in which the business of the licensee is carried on.

(4) No licence shall be issued or renewed under this Section relating to Scheduled Substances unless the person applying for or holding the licence is or has a Registered Wholesale Dealer's Licence.
Pharmacist in control of the distribution of the Scheduled Substances who is resident in Kenya.

(5) Every licence issued under this Section shall expire on the 31st day of December in the year of issue, subject to renewal.

(6) The Registrar shall keep a register of all licences issued by the Authority under this Section.

51. (1) A person carrying on a regular business in mining, agricultural or horticultural accessories shall apply to the Authority in writing on the prescribed form for a licence to deal in Scheduled Substances and any such licence, if granted shall authorize the licencee to sell only the Scheduled Substances specified therein, to persons who require them for a trade or business of mining, agricultural or horticulture.

(2) A separate licence under this Section shall be required in respect of each set of premises in which the business of the licensee is carried on.

(3) If the Authority is satisfied that it is in the public interest that a licence under this Section should be issued or renewed it may, upon payment of the prescribed fee, issue to the applicant a licence in the prescribed form, or, as the case may be, renew the licence.

(4) The Authority may refuse to issue or renew, or may revoke, a licence for any good and sufficient reason relating either to the applicant or to the licensee or to the premises in which the business is, or is proposed to be, carried on, and in case of refusal or revocation an appeal shall lie to the Cabinet Secretary, whose decision thereon shall be final.

(5) Every licence under this Section shall expire on the 31st December in the year of issue, subject to renewal.

(6) The Registrar shall maintain a register of all licences issued by the Authority under this Section.

(7) A person who sells Scheduled Substances for the purposes specified in sub-section (1) contrary to any of the provisions of this Section commits an offence and is liable upon conviction, to a fine not exceeding two hundred thousand, or to imprisonment for a term not exceeding two years, or to both.
52. (1) Subject to this Act, a person licenced under this Act to deal as a wholesaler dealer in Scheduled Substances may sell Scheduled Substances to—

(a) a person licenced under this Act as lawfully carrying on the business of a wholesale dealer in Scheduled Substances in Kenya;

(b) a person registered as lawfully carrying on the business of a pharmacist in Kenya;

(c) a person licenced under this Act as lawfully carrying on the business of a dealer in Scheduled Substances for mining, agricultural or horticultural purposes in Kenya;

(d) a qualified medical practitioner, dentist or veterinary surgeon for purposes of medical, dental or veterinary treatment respectively;

(e) the Government or a county authority or its institutions for public purposes;

(f) a hospital, dispensary or similar institution or a person or institution concerned with scientific education or research whether within or outside Kenya, where such hospital, dispensary, institution or person has been approved in that behalf by an order whether general or special, of the Cabinet Secretary: but it shall be an offence to sell Scheduled substances to any of the persons or institutions specified in paragraphs (d) and (f) unless a registered pharmacist is in direct control of the Scheduled Substances at the premises from which they are sold.

(2) Subject to this Act, an authorised seller of Scheduled Substances may sell the Scheduled Substances to any of the persons, institutions and others referred to in sub-section (1), and in addition may sell those Scheduled Substances to a person who is—

(a) in possession of the prescription of a qualified medical practitioner, dentist, or veterinary surgeon, in accordance with the prescription; or

(b) in possession of a written certificate to the effect that he may properly be supplied with the Scheduled Substances, the certificate having been issued by a person authorized by the Authority in
that behalf, a list of which persons shall be published by the Authority in the Gazette from time to time; or

(c) a person known by the seller to be a person to whom the Scheduled Substances may be properly sold.

(3) Subject to this Act, a person licenced under this Act to sell Scheduled Substances for mining, agricultural and horticultural purposes may sell Scheduled Substances in accordance with his licence.

(4) Nothing in this Section shall make it illegal for a person to sell or resell to a wholesale dealer licenced under this Act, or to an authorized seller of Scheduled Substances, stocks of Scheduled Substances which are found to be surplus to requirements, or for a person whose licence has been revoked or has expired to sell the Scheduled Substances in his possession at the time of revocation or expiry, if the sale takes place within three months after the time of revocation or expiry or such longer time as the Authority may allow.

(5) A person who sells a Scheduled Substances except in accordance with the provisions of this Section commits an offence and is liable, upon conviction, to a fine not exceeding one hundred million shillings or to imprisonment for a term not exceeding ten years or to both.

53. (1) Where a Scheduled Substance is sold in the presence of the person by whom it is to be used, the Authority may require that the seller shall not deliver it until—

(a) he has made or caused to be made an entry into the book kept for the purpose, to be called a Scheduled Substances Book, indicating in the form prescribed the date of the sale, the name and address of the purchaser and of the person, if any, by whom the certificate required under paragraph (b) of Section 53 (2) was given, the name and quantity of the Scheduled Substances sold, and the purpose for which it is stated by the purchaser to be required; and

(b) the purchaser has affixed his signature to the entry.
(2) Where a Scheduled Substances is sold in the presence of an agent or servant of the person by whom it is to be used, or where any such sale is effected by post, the following provisions apply—

(a) before the sale is completed the seller shall obtain an order in writing signed by the purchaser showing the purchaser’s name, address and occupation, the name and quantity of the Scheduled Substances to be purchased and the purpose for which it is required: provided that where a person represents that he urgently requires a Scheduled Substances for the purpose of his trade, business or profession and satisfies the seller that by reason of some emergency he is unable before delivery to furnish the order in writing, the seller may forthwith deliver the Scheduled Substances to the purchaser who shall within twenty four hours of the sale furnish the seller with the written order;

(b) before the sale is completed the seller shall satisfy himself that the signature on the order is that of the person by whom it purports to be signed, and that that person carries on the occupation stated in the order, being an occupation in which the Scheduled Substances to be purchased is properly required;

(c) the requirements of the sub-section (1) as to the making of entries in the Scheduled Substances Book shall be complied with, except that in place of the purchaser’s signature in the Scheduled Substances Book it shall be sufficient to enter in the space provided for signature the words “signed order”, together with a reference whereby the particular order may be readily identified;

(d) all signed orders and prescribed records of transactions to which this Section applies shall be retained on the premises where the sales were made, for such period as shall be prescribed

(e) if the Scheduled substance is sent by post it shall be sent by registered post or courier.

(3) A person who fails to comply with this Section commits an offence and is liable, upon conviction, to a fine
54. (1) A qualified medical practitioner, dentist or veterinary surgeon, or a member of the staff of a hospital, dispensary or similar institution who has been authorized so to do by general or special order of the Cabinet Secretary, may supply or dispense a Scheduled Substance with therapeutic value for the purpose of medical, dental or veterinary treatment, as the case may be, subject to the following provisions—

(a) the Scheduled Substance with therapeutic value shall be distinctly labelled with the name and address of the person by whom it is supplied or dispensed;

(b) the following particulars shall within twenty four hours after the Scheduled Substance with therapeutic value has been supplied or dispensed be entered in a book used regularly for the purpose, but which need not be used exclusively for that purpose, and which shall be called the Prescription Book—

(i) the date on which the Scheduled Substance with therapeutic value was supplied or dispensed;

(ii) the ingredients and the quantity supplied;

(iii) the name and address of the person to whom the Scheduled Substance with therapeutic value was supplied;

(iv) the name and address of the person by whom the prescription was given; and

(c) a registered midwife practicing domiciliary midwifery may supply or dispense a Scheduled Substance with therapeutic value in accordance with the regulations made under the Nurses Act, if he or she complies with paragraph (b) of this sub-section in relation to the supplying or dispensing of the Scheduled Substances with therapeutic value.

(2) An Authorized Seller of Scheduled Substances with therapeutic value may supply a Scheduled Substance with therapeutic value prescribed and dispensed by himself,
and in every case in which he supplies a Scheduled Substances with therapeutic value on prescription, whether the prescription has been drawn up by himself or not, shall enter the particulars in a prescription book in accordance with this Section, but shall not in respect of the supply be required to make an entry in the Scheduled Substances in the prescription book.

(3) A person to whom sub-section (1) applies who, supplies or dispenses a Scheduled Substance with therapeutic value otherwise than in compliance with these provisions commits and offence and is upon conviction, liable to a fine not exceeding one hundred thousand shillings or to imprisonment for a term not exceeding one year or both.

55. (1) It is an offence for any person to supply any Scheduled Substance unless the container of the Scheduled Substance is labelled in the prescribed manner—

(a) with the name of the Scheduled Substance;

(b) in the case of a preparation which contains a Scheduled Substance as one of the ingredients, with the prescribed particulars as to the proportion which the Scheduled Substance contained in the preparation bears to the total ingredients;

(c) with the word “Scheduled Substance” or other prescribed indication of the character of the article;

(d) if supplied on sale other than wholesale, with the name of the seller and the address of the premises on which it is sold; and

(e) if supplied otherwise than on sale, with the name and address of the supplier.

(2) The provisions of the paragraphs (a), (b) and (c) shall not apply in respect of a Scheduled Substance made up and supplied for the use of a particular person being a Scheduled Substance prescribed by reference to the needs of that person.

(3) Any person who commits an offence under this Section is upon conviction, liable to a fine not exceeding two hundred thousand shillings, or to imprisonment for a term not exceeding one year, or to both.
56. A person exposing or causing to be exposed for sale any Scheduled Substance in or by means of an automatic machine commits an offence and shall be liable to a fine not exceeding five hundred thousand shillings, or to imprisonment for a term not exceeding three year, or to both.

57. (1) This shall be permitted as long as the supply of the medicine conforms with all requirements for the particular medicine in terms of its scheduling status and any other requirements as may be specified in Regulations pertaining to this type of supply. In the case of a Prescription-only medicine, the required prescription shall have been obtained as a result of at least one physical interaction between an authorised practitioner and the patient.

(2) Any online pharmacy or other electronic source of supply of medicines which does not comply with these provisions shall be guilty of an offence and shall be liable to a fine not exceeding two hundred thousand shillings, or to imprisonment for a term not exceeding one year, or to both.

PART VI—MANUFACTURE OF MEDICINAL SUBSTANCES

58. (1) A person shall not manufacture any medicinal substance unless the person has been granted a manufacturing licence by the Authority.

(2) Each manufacturing licence shall expire on the 31st December of every year and the renewal shall be subject to the compliance with conditions prescribed by the Authority.

(3) A person shall not manufacture any medicinal substance for sale unless the person has applied for and obtained a licence from the Authority in respect of each substance intended to be manufactured.

(4) Any person who intends to manufacture a medicinal substance shall make an application in the prescribed form for the licensing of the premises and the application shall be accompanied by the prescribed fee.

(5) In issuing a licence, the Authority may prescribe any licencing conditions that it considers necessary.
59. A person who is granted a manufacturing licence under Section 59 shall comply with the good manufacturing practices prescribed by the Authority.

PART VII—THERAPEUTIC COSMETICS

60. (1) A person shall not sell any therapeutic cosmetic that—

(a) contains any substance that may cause injury to the health of the user when the therapeutic cosmetic is used—

(i) according to the directions on the label of or accompanying such therapeutic cosmetic; or

(ii) for such purposes and by such methods of use as are customary or usual thereof; or

(iii) consists in whole or in part of any filthy, disgusting, rotten, decomposed or diseased substance or of any injurious foreign matter; or

(iv) was prepared, preserved, packed or stored under insanitary conditions,

(2) A person who contravenes subsection (1) commits an offence.

61. If a standard has been prescribed for a therapeutic cosmetic, any person who labels, packages, sells or advertises any article in such a manner that it is likely to be mistaken for a therapeutic cosmetic of the prescribed standard commits an offence.

62. Any person who sells, prepares, preserves, packages, conveys, stores or displays for sale any therapeutic cosmetic under insanitary conditions commits an offence.

63. Any therapeutic cosmetic which either contains a Scheduled Substance or claims to have a therapeutic effect or value shall be treated as a medicine.

64. The Registrar shall keep a register of all therapeutic cosmetics registered under section 64 which shall be called a therapeutic cosmetics register.

65. (1) The Authority, in the public interest, may prohibit any ingredient contained in therapeutic cosmetics by notice in the Gazette.

(2) Except as otherwise provided in the regulations, a
cosmetic shall not contain any prohibited ingredients.

(3) Any person who manufactures or knowingly sells a therapeutic cosmetic which contains a prohibited ingredient commits an offence.

PART VIII — MEDICAL DEVICES

66. (1) The Registrar shall keep in the prescribed form a human medical devices register and a veterinary medical devices register, to be known as the medical devices registers in which the registrar shall register all medical devices, the registration of which has been approved by the Authority.

(2) The Register under subsection (1) shall contain all such particulars in regard to such medical devices and the holder of the certificate of registration in respect of such medical devices as are required by this Act or any other law to be entered therein.

67. (1) A person shall not sell any medical device—

(a) that is not registered by the Authority

(b) that is adulterated;

(c) substandard;

(d) defective; or

(e) which fails to comply in any way with specifications of this Act or any other law.

(2) Any person who sells any medical device that, when used according to directions on the label or contained in a separate document delivered with the medical device or under such conditions as are customary or usual, may cause injury to the health of the purchaser or its user commits an offence.

68. Any person who labels, packages, treats, processes, sells or advertises any medical device in contravention of this Act or any other law, or in a manner that is false, misleading or deceptive as regards its character, value, composition, merit or safety, commits an offence.

69. (1) Where a standard has been prescribed for a medical device, any person who labels, packages, sells or advertises any article in such a manner that it is likely to be mistaken for that medical device commits an offence unless
the article complies with the prescribed standard.

(2) The Authority may issue standards to ensure that medical device are designed and produced in a way that ensures that the exposure of a patient, the user, or any other person, to radiation is minimised, having regard to the levels of radiation required to enable the device to perform its therapeutic and diagnostic functions and the intended purpose of the device.

70. (1) A person who sells, manufactures, packages, stores or conveys for sale any medical device under insanitary conditions commits an offence.

(2) A person who knowingly sells a medical device that has a measuring function that does not provides accurate, precise and stable measurements within the limits indicated by the manufacturer and having regard to the intended purpose of the device commits an offence.

(3) A person sells or supplies unapproved medical devices commits an offence.

PART IX—BLOOD AND BLOOD PRODUCTS

71. (1) A person may not undertake any of the following activities without a licence from the Authority—

(a) the collection and testing of blood or blood components, whatever their intended purpose; and

(b) the processing, storage and distribution of blood and blood components products when they are intended to be used for transfusion.

(2) Paragraph (1) shall not apply to—

(a) the storage and distribution of, and the performance of compatibility tests on, blood and blood components exclusively for use within hospital facilities, including transfusion activities where such activities are performed by a hospital blood bank; or

(b) any person carrying out any of the activities referred to in paragraph (1), where that person carries out that activity on behalf of, and pursuant to a contractual arrangement with—

(i) a blood establishment which is authorised

Licences to deal in blood and blood products.

Offences relating to medical devices.
under act or any other law to carry out the activity in question; or

(ii) a person responsible for management of a hospital blood bank.

(3) A person who contravenes subsection (1) commits an offence.

(4) The Authority may licence a person who intends to carry out any of the activities referred to in subsection (1).

(5) An application for a licence under subsection (1) shall be made to the authority in the prescribed form.

(6) The Authority shall prescribe through regulations—

(a) the procedure for applying for a licence under this section;
(b) the licensing requirements;
(c) the application fees to be paid by an applicant;
(d) any other condition or requirement for licencing;
(e) the requirements of a blood establishment;
(f) labeling of blood and blood components;
(g) requirements for hospital blood banks.

(7) A person licenced under subsection (4) of section (72) shall keep a record of—

(a) donors and recipients;
(b) results from laboratory tests carried out on blood and blood products;
(c) every unit of blood and blood components collected;
(d) the activities relating to collection, testing, processing, labeling, Documenting, distribution, storage, and

(e) use of blood and blood products;

(f) The destruction of every unit of blood and the reasons for the destruction; and

(g) any other information that the Authority may prescribe.

(8) The information under paragraph 1 shall be kept
confidential and any person who contravenes this subsection commits an offence.

(9) The Authority may suspend or revoke the authorisation of a blood establishment on one or more of the following grounds—

(a) that the blood establishment has failed, in any material respect, to comply with the requirements of this Act or any other law;

(b) that the collection, testing, processing, storage or distribution of blood or blood components by the establishment cannot be carried out safely;

(c) that any blood or blood components cannot be supplied to hospital blood banks in such a state that they could be safely administered for transfusion;

(d) that the information given by the blood establishment pursuant in applying for a licence was false or incomplete in any material respect or

(e) any other reason that the authority considers appropriate.

(10) Before suspending or revoking the authorisation of a blood establishment, the Authority shall serve a notice on the blood establishment stating that the authority intends to suspend or revoke its authorisation with effect from the date specified in the notice, which date shall be not less than ten days from the date on which the notice is served.

(11) Where the authority considers that it is necessary in the interests of safety, it may, by a notice served on a blood establishment, suspend or revoke its authorisation with immediate effect.

(12) Where—

(a) the blood establishment has failed in any material respect, to comply with the requirements of this Act; or

(b) the information given by the blood establishment pursuant in an application of a licence was false or incomplete in any material respect, and the Authority considers that the failure in question is not sufficiently serious to warrant suspension or revocation of the authorisation of the blood
establishment in the first instance, the may serve a notice on the responsible person of the blood establishment in accordance with paragraph (12).

(13) A notice served under this paragraph shall—

(a) identify the requirements of the regulations of which the blood establishment is in breach or, in the case of false and incomplete information, the further information which is required;

(b) identify the action which the blood establishment is required to take; and

(c) give the timescale within which the blood establishment shall take the action identified in sub-paragraph (b).

(14) If the blood establishment fails to comply with the requirements set out in the notice within the specified timescale, the Authority may, by a notice served on the blood establishment, suspend or revoke the authorisation of the blood establishment.

(15) A suspension or revocation pursuant to paragraph (6) shall take effect—

(a) in a case where the Authority considers that it is necessary in the interests of safety, immediately; or

(b) in all other cases, from a date specified in the notice.

(16) Any suspension pursuant to this Act shall be for such period as the Authority shall consider necessary having regard to the reasons for the suspension.

(17) The suspension or revocation of an authorisation may be total, or may be limited to a particular activity or to one or more activities carried out at a particular site or sites, or to a particular blood component.

72. (1) Collection of blood in exchange for payment by the health care establishments under Article 15 is permitted in the following cases—

(a) In cases of emergency and in the absence of available quantities of the necessary blood group at the corresponding Centre for Transfusion.
Haematology;

(b) For manufacturing of vaccines, sera, and immunoglobulins;

(c) For research and diagnostic medical purposes.

(2) Despite subsection (1), a health care establishments may cover the direct costs of blood donors, and incite them with symbolic gifts or by means of another way that is compatible with the principle of voluntariness of blood donations.

(3) There is established a blood and blood products register.

(4) The blood and blood products register shall contain—

(a) the health care establishments that implement activities relating to collection, testing, processing, storage, and distribution of blood and blood components; and

(b) the serious incidents and the serious adverse reactions associated with collection and use of blood and blood components.

73. (1) Any person who carries out collection, testing, processing, transfusion, and storage of blood or blood components shall immediately notify the Authority of any serious incidents or adverse reactions that have occurred with respect to blood and blood components.

(2) The Authority through authorized persons shall analyze and summarize the information regarding the serious incidents and the serious adverse reactions and shall undertake measures for their mitigation and prevention.

(3) Blood and blood components that do not comply with the standards prescribed by the authority shall be withdrawn from use and shall be destroyed or delivered for educational or research needs with the permission of the Authority under such terms and conditions as may be determined by the Authority.

(5) A person who contravenes subsection (3) commits an offence.
PART X—TOBACCO PRODUCTS

74. (1) A person shall not manufacture a tobacco product that does not conform with the standards established under this Act or any other law.

(2) A person who contravenes subsection (1) commits an offence.

75. Every manufacturer of a tobacco product shall submit to the Authority, in the prescribed manner and within the prescribed time, information that is required by the Authority about the manufacturer's tobacco products, their emissions and any research and development related to tobacco products and their emissions, whether the tobacco products are for sale or not.

76. (1) The Authority may make regulations—

(a) establishing standards for tobacco products, including prescribing the amounts of substances that may be contained in the product or its emissions;

(b) respecting test methods, including methods to assess conformity with the standards;

(c) prescribing information that manufacturers must submit to the Authority about tobacco products and their emissions, including sales data and information on market research, product composition, ingredients, health effects, hazardous properties and brand elements;

(d) prescribing information that manufacturers must submit to the Authority about research and development related to tobacco products and their emissions, including information on market research, product composition, ingredients, health effects, hazardous properties and brand elements;

(e) respecting the prohibition of any additives, including providing for the suspension of the manufacture or sale of a tobacco product; and

(f) prescribing the means, including electronic means, by which the information referred to in paragraphs (c) to may be submitted to the Authority.
PART XI—NATIONAL QUALITY CONTROL LABORATORY

77. (1) There is established a National Quality Control Laboratory of the Authority which shall be used as a facility for—

(a) the examination and testing of the drugs and any material or substance from or with which and the manner in which drugs may be manufactured, processed or treated and ensuring the quality control of drugs and medicinal substances;

(b) performing chemical, biological, biochemical, physiological and pharmacological analysis and other pharmaceutical evaluation;

(c) conduct research and training; and testing the quality of locally manufactured and imported medicines or medicinal substances, therapeutic foods, medical devices or therapeutic cosmetics on behalf of the Authority, with a view to determining whether such drugs or medicinal substances comply with this Act or rules made hereunder.; and

(d) do such other function as shall be determined by the Authority.

78. (1) A certificate of analysis shall be issued and signed by the Director General for every analysis done by the National Quality Control Laboratory.

(2) The certificate of analysis issued under subsection (1) shall be in the prescribed form.

PART XII—ADVERTISEMENTS AND LABELLING

79. (1) Subject to the provisions of this Act, no person shall advertise any health product and technology except with the written permission of the Authority.

(2) Applications for the advertisement of any health product and technology shall be made to the Authority in the prescribed form and shall be accompanied by the prescribed fee.

80. (1) Subject to this Act, a person shall not take part in the preparation or publication of an advertisement
referring to a medicine, drug, appliance or article of any description in terms which are calculated to imply that the medicine, drug, appliance or article may be effective for any of the purposes specified in the Sixth Schedule under this Act.

(2) In proceedings for the contravention of subsection (1), it shall be a defence for the person charged to prove that the advertisement to which the proceedings relate was published only so far as was reasonably necessary to bring it to the notice of one or more persons of the following classes—

(a) members of Parliament;
(b) members of the board of a hospital;
(c) duly qualified medical practitioners, dentists and veterinary surgeons;
(d) registered pharmacists, Authorized Sellers of Scheduled Substances and licenced wholesale dealers; or
(e) persons carrying on business which includes the sale or supply of surgical appliances, or that the advertisement was so published in connexion with an application for a patent submitted to the appropriate authority so far only as was requisite for the purpose of the application.

(3) The Cabinet Secretary may from time to time, by notice in the Gazette, amend or vary the Schedule.

81. Subject to this Act, a person shall not take part in the publication of an advertisement referring to a medicine, drug, appliance or article of any description, in terms which are calculated to lead to the use of the drug, appliance or article for procuring the miscarriage of women.

82. (1) Subject to this Act—

(a) a person shall not take part in the publication of an advertisement referring to a health product or technology or similar article in terms which in the opinion of the Authority are considered to be extravagant, false or misleading and to bear little or no relation to the pharmacological properties etc. etc.
and action of the ingredients or the component or correct use of the article.

(b) a person shall not in an advertisement claim that the therapeutic efficacy of a health product or technology or use of an article is other than that for which the drug has been registered by the Board in terms of this Act or state or suggest that such health product or technology should be used for a purpose, under a circumstance, or in a manner, other than that for which it is registered by the Authority.

83. (1) In proceedings for contravention of any of the provisions of Sections 83 and 84—

(a) that an advertisement was published referring to a drug, appliance or article of any description, in terms calculated to lead to the use of the drug, appliance or article—

(i) in the case of contravention of Section 83, for the treatment of any of the human ailments referred to in subsection (1) of that Section; or

(ii) in the case of a contravention of Section 84, for procuring the miscarriage of women; and

(b) that the advertisement also referred to the medicine, drug, appliance or article in terms calculated to indicate that it was manufactured, produced, imported, sold or offered for sale by the person charged,

unless the contrary is proved, it shall be presumed for the purpose of those proceedings that that person took part in the publication of the advertisement, but without prejudice to the liability of any other person.

(3) In proceedings for contravention of any of the provisions of Sections 82, 83, 84 and 85, it shall be a defence for the person charged to prove—

(a) that the advertisements to which the proceedings relate was published in such circumstances that he did not know and had no reason to believe that he was taking part in the publication; or
(b) that the advertisement was published only in a publication of a technical character intended for circulation only amongst persons of the following classes, or of one or some of them—

(i) qualified medical practitioners, dentists and veterinary surgeons;

(ii) registered pharmacists and authorized sellers of Scheduled Substances;

(iii) persons undergoing training with a view to becoming qualified medical practitioners, dentists or veterinary surgeons, or registered pharmacists; or

(i) persons carrying on business which includes the sale or supply of surgical appliances.

84. (1) Subject to this Act, a person shall not sell by retail an article consisting of or comprising a substance recommended as a medicine unless there is written so as to be clearly legible on the article or on a label affixed thereto, or if the article is sold or supplied in more than one container, on the inner container or on a label affixed thereto—

(a) the appropriate designation of the substance so recommended or of each of the active constituents, or of each of the ingredients from which it has been compounded; and

(b) in a case where the appropriate designation of each of the active constituents or ingredients is written, the appropriate quantitative particulars of the constituents or ingredients; provided that this subsection shall not apply to an article made up and supplied for the use of a particular person, being an article prescribed by reference to the needs of that person.

(2) In sub-section (1)—

"appropriate designation", in relation to a substance, constituent or ingredient, means—

(a) in a case where the substance, constituent or ingredient is a scheduled substance included in
the schedules under this Act, the name with which the container of the scheduled substance is for the time being required to be labelled in accordance with this Act or regulations made under this Act; or

(b) in a case where the substance, constituent or ingredient is not such a scheduled substance and is described in any of the monographs contained in the edition of the British Pharmacopoeia or the British Pharmaceutical Codex or the International Pharmacopoeia or the British Veterinary Codex which was last published before the date on which the article was sold or supplied, the description set out at the head of that monograph;

(c) in a case where the substance, constituent or ingredient is not such a scheduled substance and is not so described, the accepted scientific name, or other name descriptive of the true nature of the substance, constituent or ingredient, and in all cases the appropriate name of the substance shall be written in English;

"appropriate quantitative particulars", in relation to the active constituents or the ingredients of a substance, means—

(a) the approximate percentage of each of those constituents or ingredients contained in the substance or the approximate quantity of each of those constituents or ingredients contained in the article sold or supplied or contained in a defined quantity of the article; or

(b) in the case where the article consists of or comprises a number of separate portions of the substance, either the approximate percentage or quantity or the approximate quantity of each of the constituents or ingredients contained in each portion;

"container" includes a wrapper.

(3) If a person sells or supplies an article in contravention of this Section, subject to this Act, the person
commits an offence and upon conviction, is liable—

(a) in the case of a first conviction, to a fine not exceeding two hundred thousand shillings or to an imprisonment term not exceeding two years or to both;

(b) in the case of a subsequent conviction, to a fine of at least three hundred thousand shillings or to imprisonment to a term not exceeding three years or to both.

85. (1) It shall be a defence for a person charged with selling or supplying, in contravention of Section 80, an article consisting of or comprising a substance recommended as a medicine to prove—

(a) that the person did not know and had no reason to believe, that the article consisted of or comprised such a substance; or

(b) that, in relation to the matter in respect of which the person is charged, the person acted in the course of the person’s employment as a servant or agent of another person on the instruction of the person’s employer or of some other specified person.

(2) In proceedings for contravention of Section 80 a document purporting to be a certificate signed by a public analyst or by an officer authorized in writing by the Cabinet Secretary to perform such analysis, and stating the result of an analysis made by the person, shall be admissible as evidence of the matters stated therein, but a party to the proceedings may require the person by whom the analysis was made to be called as a witness.

86. Where a person is charged with an offence under this Act by a reason of the person having sold or been in possession of a container as containing an article, and the container seems to have been packed by the manufacturer of the contents and to be intact, the container shall be presumed to contain articles of the description specified on the label, until the contrary is proved.

87. An appeal under any Section shall be in writing and shall be lodged within 30 days after the date of the act appealed against.
PART XIII—ADMINISTRATION AND ENFORCEMENT

88. (1) The Cabinet Secretary, on the recommendation of the Authority may, by order, prohibit or control the manufacture, sale, advertisement or possession of any secret, patent, proprietary or homoeopathic medicine, preparation or medical device.

(2) A person who contravenes an order made by the Cabinet Secretary under subsection (1) commits an offence.

89. (1) The Authority may authorise a person, in writing, to supply a specified quantity of a particular health product or technology, which is subject to registration under this Act, but is not registered, during a specified period and to a specified person or institution.

(2) A health product or technology supplied under the authority granted in terms of this section may be used for such purposes, in such manner and during such period, as the Authority may determine in writing.

(3) Subsequently, if effect is not given to a determination made in terms of this section, or if the Authority is of the opinion that the risks of supplying a specified quantity of a particular medicine or medical device product in terms of this section, outweigh the potential benefits, the Authority may at any time, in writing, withdraw any such authority granted.

90. (1) A drug, article or document seized under the provisions of this Act may be retained for a period not exceeding one month or if within that period proceedings are commenced for an offence under this Act in respect of that drug, article or document, until the final determination of those proceedings.

(2) Where a magistrate is satisfied that any such drug or article is of a perishable nature or that by reason of the fact that the market for the drug or article is seasonal, or for any other reason, delay in disposing the drug or article would unduly prejudice the owner, the magistrate may authorize the sale or other disposal of the drug or article.

(3) where proceedings are taken for an offence under this Act or any rules there under the court by or before which the alleged offender is tried may make such order as
to the forfeiture or other disposal of any drug or article in respect of which such offence was committed as the court shall see fit.

(4) In this Section references to a drug or article shall be construed as including the proceeds of a sale effected in accordance with the provisions of subsection (2).

91. The Authority may—
(a) by notice in writing, require a person, who manufactures, supplies, administers or prescribes a health product or technology, or on whose direction a health product or technology is manufactured, supplied or administered, to furnish the Authority, within a period specified in that notice, with information, which is in that person's possession or which that person is in a position to obtain with respect to that health product or technology.
(b) if requested by a person to whom a notice under this section is addressed, extend the period specified in that notice.

92. (1) An authorized or licenced seller of any food, scheduled substance health product or technology shall, on the demand of a regulatory officer, produce for inspection his certificate of registration or his licence as the case may be.
(2) All books kept by any seller of scheduled substance a food, health product or technology, medical practitioner, dentist or veterinary surgeon, or by a hospital, dispensary or similar institution, in accordance with the provisions of this Act or any rules there under, shall be open for inspection by a regulatory officer at all reasonable times.

93. A person who obstructs or hinders a regulatory officer in the lawful exercise of the powers conferred by this Act commits an offence.

94. (1) An act which if done by an individual would be an offence under this Act or by any rules thereunder shall, if done by a body corporate, be an offence by every director, secretary and manager unless it is proved that the offence was committed without individual's consent or
connivance and that the individual exercised all such
diligence to prevent the commission of the offence as the
individual ought to have exercised having regard to the
nature of the individual's functions in that capacity and to
all the circumstances.

(2) If an offence against this Act or any rules
thereunder has been committed by a partner in a firm, every
person who at the time of the commission of the offence
was a partner in that firm, or was purporting to act in that
capacity, shall be deemed to be guilty of that offence unless
the person proves that the offence was done without the
person's consent or connivance and that the person
exercised all such diligence to prevent the commission of
the offence as the person ought to have exercised having
regard to the nature of the person's functions in that
capacity and to all the circumstances.

95. (1) If—

(a) a body corporate has been convicted of an
offence under this Act or any other rules
thereunder; or

(b) a member of the Authority or an officer of a body
corporate, or a person employed by a body
corporate in carrying on a business has been
convicted of any such criminal offence, or been
guilty of misconduct which in the opinion of the
Authority renders the person, or would if the
person were a registered pharmacist render the
person, unfit to be on the register, then, whether
the body corporate was or was not an Authorized
Seller of scheduled substance health product or
technology at the time when the offence or the
misconduct was committed, the Authority may
inquire into the case and may, subject to this Act
direct—

(i) that the body corporate shall, in a case
where it is an Authorized Seller of
scheduled substance health product or
technology, cease to be a seller and, in any
case, be disqualified for such period as may
be specified in the direction from being an
authorized seller of scheduled substance
health product technology; or
(ii) that all or any other of the premises of the body corporate shall, in a case where they are registered in the register of premises kept in pursuance under this Act, be removed from that register and in any case be disqualified for such period as may be specified in the directions from being registered therein.

(2) A body corporate may appeal to the Cabinet Secretary against a direction given under this Section.

96. (1) Where any regulations made under this Act or under the Public Health Act prohibits or restricts the addition of any preservative or other ingredient or material to any food, the addition of such preservative, ingredient or material, if made in contravention of the regulations, shall, for the purposes of this Act, be deemed to render the food injurious to health.

(2) Where any regulations made under this Act or under the Public Health Act prescribes the composition of any article of food intended for sale, or prohibit or restrict the addition of any preservative or other ingredient or material to any such article, the purchaser of such article shall, unless the contrary is proved, be deemed for the purposes of this Section to have demanded an article complying with the provisions of the regulations as regards the presence or amount of any constituent, ingredient or material specified in the regulations.

97. A regulatory officer may, for the purposes of this Act, inspect any animal intended for slaughter and may seize and examine any meat which the regulatory officer considers to be unfit for consumption.

98. (1) A regulatory officer may, at any hour reasonable for the proper performance of duty—

(a) enter any premises where the regulatory officer believes any article to which this Act or any regulations made thereunder apply is prepared, preserved, packaged, stored or conveyed, examine any such article and take samples, and examine anything that the regulatory officer believes is used or capable of being used for such preparation, preservation, packaging or storing or conveying;
(b) stop or search or detain any aircraft, ship or vehicle in which the regulatory officer believes that any article subject to the provisions of this Act is being conveyed and to examine any such article and take samples for the purposes of this Act;

(c) open and examine any receptacle or package which the regulatory officer believes contains any article to which this Act or any regulations made thereunder apply;

(d) examine any books, documents, or other records found in any place mentioned in paragraph (a) of sub-Section (1) that the regulatory officer believes contain any information relevant to the enforcement of this Act with respect to any article to which this Act or any regulations made apply and make copies or take extracts;

(e) seize and detain for such time as may be necessary any article by means of or in relation to which he believes any provision of this Act or any regulations made thereunder has been contravened.

(2) A regulatory officer acting under this Section shall, produce his authority.

(3) Any owner, occupier or person in charge of any premises entered by a regulatory officer pursuant to paragraph (a) of subsection (1), or any person found therein, who does not give to the regulatory office all reasonable assistance the person’s power and furnish the regulatory officer with such information as the regulatory officer may reasonably require, shall be guilty of an offence.

(4) Any person who obstructs or impedes any regulatory officer in the course of the regulatory officer’s duties or by any gratuity, bribe, promise, or other inducement prevents, or attempts to prevent the due execution by the regulatory officer of the regulatory officer’s duty under this Act or any regulations made thereunder commits of an offence.

(5) Any person who knowingly makes any false or misleading statement either verbally or in writing to any regulatory officer commits an offence.
(6) A regulatory officer shall release any article seized by the regulatory officer under this Act when the regulatory officer is satisfied that all the provisions of this Act and any regulations made thereunder with respect thereto have been complied with.

(7) Where a regulatory officer has seized an article under this Act and the owner thereof or the person in whose possession the article was at the time of seizure consents to the destruction thereof, the article may be destroyed or otherwise disposed of as the regulatory officer may direct.

(8) Where a person has been convicted of an offence under this Act or any regulations made thereunder, the court may order that any article by means of or in relation to which the offence was committed or anything of a similar nature belonging to or in the possession of the convicted person or found with such article, be forfeited, and upon such order being made such articles and things may be disposed of as the court may direct.

(9) Where any article has been seized under the provisions of paragraph (e) of subsection (1) and the owner thereof has been convicted of an offence under this Act, the article may be destroyed or otherwise disposed of as the regulatory officer may direct.

(10) Any article seized under this Act may at the option of a regulatory officer be kept or stored in the premises where it was seized or may at the direction of a regulatory officer be removed to any other proper place; and any person who removes, alters or interferes in any way with articles seized under this Act without the authority of a regulatory officer shall be guilty of an offence.

(11) A regulatory officer may submit any seized article or any sample therefrom to the National Quality Control Laboratory for analysis or examination; and a public analyst shall as soon as practicable analyse or examine any sample sent to to the public analyst in pursuance of this Act and shall give the regulatory officer a certificate specifying the result of the analysis or examination, and such certificate shall be in such form as may be prescribed by the Authority.

(12) In this section, "premises" includes a street, open space, place of public resort, or bicycle or other vehicle
utilized for the preparation, preservation, packaging, storage or conveyance of any article.

(13) In performing any of the functions under this Act, the regulatory officer, as the situation may require, may be accompanied and assisted by a Police Officer.

(14) The procedure to be followed by regulatory officer in obtaining, transmitting for analysis or examination or otherwise dealing with any sample, shall be prescribed by this Act or any other law.

99. The Director of Medical Services, or , in relation to any matter appearing to affect the general interests of the consumer, the Director of Veterinary Services , , in relation to any matter appearing to affect the general interests of animal husbandry in Kenya, and the Director of Agriculture in relation to any matter appearing to affect the general interests of agriculture in Kenya, and any other person authorized in writing by the Cabinet Secretary so to do, may direct a public officer to procure for analysis samples of any food, health product or technology, and thereupon that officer shall have all the powers of a regulatory officer under this Act and this Act shall apply as if the officer were a regulatory officer.

100. (1) It shall be the duty of every county authority to exercise such powers with which it is vested as may be, in its special circumstances, reasonably practicable so as to provide proper safeguards for the sale of food, scheduled substance and health product and technologies in a pure and genuine condition, and in particular to direct its officers to procure samples for analysis.

101. (1) The Cabinet Secretary may direct any person who at the date of the direction or at any subsequent time carries on a business which includes the production, importation or use of substances or articles of any class to which this Act applies to furnish to the cabinet secretary, within such time as may be specified in such direction, such particulars as may be so specified of the composition and use of any such substance or article sold or for sale in the course of that business or used in the preparation of any article.

(2) Without prejudice to the generality of subsection (1), a direction made thereunder may require the following particulars to be furnished in respect of any substance—
particulars of the composition and chemical formula of a substance;

(a) particulars of the manner in which a substance is used or proposed to be used in the preparation of food;

(b) particulars of any investigations carried out by or to the knowledge of the person carrying on the business in question, for the purpose of determining whether and to what extent the substance, or any product formed when the substance is used as aforesaid, is injurious to, or in any other way affects health;

(c) particulars of any investigations or inquiries carried out by or to the knowledge of the person carrying on the business in question for the purpose of determining the cumulative effect on the health of a person consuming the substance in ordinary quantities.

(3) No particulars furnished in accordance with a direction under this section, and no information relating to any individual business obtained by means of such particulars shall, without the previous consent in writing of the person carrying on the business in question, be disclosed except—

(a) in accordance with regulations made by the Authority, so far as may be necessary for the purposes of this Act; and

(b) for the purposes of any proceedings for an offence against the order or any report of those proceedings, and any person who discloses any such particulars or information in contravention of this subsection shall be guilty of an offence.

102. On the conviction of any person for any offence under this Act or any regulations made under the Act, the court may, in addition to or in lieu of any other penalty which it may lawfully impose, cancel any licence issued under this Act, or any regulations made under the Act, to such person.

103. (1) A Regulatory Officer may take out proceedings for an offence under this Act or the regulations before any magistrate having jurisdiction in the place where
any article sold was actually delivered to the purchaser or where the sample was taken.

(2) In any proceedings under this Act, the contents of any container appearing to be intact and in the original state of packing by the manufacturer thereof shall be deemed, unless the contrary is proved, to be an article of the description specified on the label.

104. (1) A person who is guilty of an offence under this Act for which no special penalty is provided shall be liable—

(a) in the case of a first offence, to a fine not exceeding five hundred thousand shillings or to imprisonment for a term not exceeding one year, or to both such fine and imprisonment;

(b) in the case of a subsequent offence, to a fine not exceeding seven hundred thousand shillings or to imprisonment for a term not exceeding five years, or to both such fine and imprisonment.

(2) In any prosecution under this Act the summons shall state the particulars of the offence or offences alleged and also the name of the prosecutor and shall not be made returnable in less than fourteen days from the date on which it is served.

105. In any proceedings under this Act—

(a) a certificate of analysis purporting to be signed by a National Quality Control Laboratory shall be accepted as prima facie evidence of the facts stated:

(b) despite (a) above—

(i) the party against whom it is produced may require the attendance of the public analyst for the purposes of cross-examination; and

(ii) no such certificate of a public analyst shall be received in evidence unless the party intending to produce it has, before the trial given to the party against whom it is intended to be produced, reasonable notice of such intention together with a copy of the certificate;
(c) evidence that a package containing any article to which this Act or any regulations made thereunder apply bore a name, address or registered mark of the person by whom it was manufactured or packed shall be prima facie evidence that such article was manufactured or packed, as the case may be, by that person;

(d) any substance commonly used for human consumption shall, if sold or offered, exposed or kept for sale, be presumed, until the contrary is proved, to have been sold or, as the case may be, to have been or to be intended for sale for human consumption;

(e) any substance commonly used for human consumption which is found on premises used for the preparation, storage, or sale of that substance and any substance commonly used in the manufacture of products for human consumption which is found on premises used for the preparation, storage or sale of those products, shall be presumed, until the contrary is proved, to be intended for sale, or for manufacturing products for sale, for human consumption; or

(f) any substance capable of being used in the composition or preparation of any substance commonly used for human consumption which is found on premises on which that substance is prepared shall, until the contrary is proved, be presumed to be intended for such use.

PART XIV—FINANCIAL PROVISIONS

106. The funds of the Authority shall consist of—

(a) Appropriations and budgetary allocations from the Consolidated Funds;

(b) such monies or assets as may accrue to the Authority in the course of the exercise of its powers or the performance of its functions under this Act;

(c) gifts, grants or donations as may be given to the Authority;

(d) monies that may be borrowed by the Board for
the discharge of the functions of the Authority;
and

(e) monies from any other source

107. The financial year of the Authority shall be the period of twelve months ending on the thirtieth day of June in each year.

108. (1) At least three months before the commencement of each financial year, the Board shall cause to be prepared estimates of the revenue and expenditure of the Authority of that year.

(2) The annual estimates shall make provisions for all estimated expenditure of the Authority for the financial year concerned and in particular, shall provide for—

(a) the payment of the salaries, allowances and other charges in respect of the staff of the Authority;

(b) the payment of pensions, gratuities and other charges in respect of benefits which are payable out of the funds of the Authority;

(c) the acquisition and maintenance of the buildings and grounds of the Authority;

(d) the funding of training, research and development activities of the Authority;

(e) the proper maintenance, repair and replacement of any installation and of the equipment and other movable property of the Authority;

(f) the creation of such funds to meet future or contingent liabilities in respect of benefits, insurance or replacement of buildings or installation or equipment and in respect of such other matters as the Authority may think fit.

(3) The annual estimates shall be approved by the Authority before the commencement of the financial year to which they relate, and shall be submitted to the Cabinet Secretary for approval and after the Cabinet Secretary has given approval, the Authority shall not increase any sum provided in the estimates without written consent of the Cabinet Secretary.

(4) No expenditure shall be incurred for the purposes of the Authority except in accordance with the annual
estimates approved under subsection (3), or in pursuance of
an authorization of the Authority given with the prior
approval of the Cabinet Secretary.

109. (1) The Authority shall cause to be kept all
proper books and records of account of the income,
expenditure, assets and liabilities of the Authority.

(2) The Cabinet Secretary for the time being
responsible for finance may prescribe the form of any book
required to be kept under subsection (1) and unless a form
has been prescribed, a form suitable for the purpose shall
be used.

(3) Within a period of three months after the end of
each financial year, the Authority shall submit to the Kenya
National Audit Office the accounts of the Authority in
respect of that year together with—

(a) a statement of the income and expenditure of the
Authority during the financial year; and

(b) a statement of the assets and liabilities of the
Authority on the last day of that financial year.

(4) The accounts of the Authority shall be audited and
reported upon by the Kenya National Audit Office.

110. (1) The Authority may invest any of its funds in
securities in which for the time being trustees
may by law invest trust funds or in any other
securities which the Treasury may, from time to
time approve.

(2) The Authority may place on deposit with such
bank or banks or financial institutions as it may
determine, any moneys not immediately required
of the purposes of the Authority.

111. (1) The Authority shall cause an annual report to
be prepared for each financial year.

(2) The Authority shall submit the annual report to the
Cabinet Secretary within three months after the end of
the year to which it relates.

(3) The annual report shall contain, in respect of the
year to which it relates—

(a) the financial statements of the Authority;
(b) a description of the activities of the Authority;

(c) such other statistical information as the Authority considers appropriate relating to the work of the Authority;

(d) any other information relating to the functions that the Authority considers necessary.

(4) The Cabinet Secretary shall, within thirty days, after receiving the annual report, transmit it to the National Assembly.

112. The Authority may, at any time, submit a special report to the National Assembly with respect to any aspect of the functions of the Authority which the Authority considers should, in the national interest, be brought to the attention of the National Assembly because it affects a wide cross-section of the populace and there could be disastrous consequences if a report thereon is not brought to the attention of the National Assembly.

PART XV — MISCELLANEOUS PROVISIONS

113. (1) The Authority shall make regulations for the better carrying out of the provisions of this Act.

(2) Without prejudice to the generality of sub-section (1), the authority may make regulations—

(a) setting out the procedure and criteria for determining and declaring that any food is adulterated;

(b) with respect to—

(i) the labelling and packing and the offering, exposing and advertising for sale of food, drugs, health products and technologies;

(ii) the size, dimensions and other specifications of packages of food, health products and technologies;

(iii) the sale or the conditions of sale of any food, drug, health products and technologies; and

(iv) the use of any substance as an ingredient in any food, health products and technologies;

(c) to prevent the consumer or purchaser from being
deceived or misled as to its quality, quantity, character, value, composition, effect, merit or safety of any food, health product and technology or to prevent injury to the health of the consumer or purchaser;

(d) prescribing the type and level of food additives;

(e) the treatment, processing and manufacture and distribution of food;

(f) prescribing standards of composition, strength, potency, purity, quality or other property of any food, health products and technologies;

(g) respecting the importation or exportation of food, health product and technologies in order to ensure compliance with this Act or any other law;

(h) respecting the method of preparation, preserving, packing, storing, conveying, distribution and testing of any food, health products or technologies;

(i) respecting the carriage of goods subject to the provisions of this Act, including the licensing of vehicles used in such carriage;

(j) requiring persons who sell food, health products and technologies to maintain such books and records as the authority considers necessary for the proper enforcement and administration of this Act and any other law;

(k) requiring manufacturers of any drugs or scheduled

(l) substances to submit test portions of any batch of such drugs or scheduled substances;

(m) providing for the analysis of food, health products and technologies for the purposes of this Act or for any other purpose;

(n) prescribing a tariff of fees to be paid for such analysis and for prescribing methods of analysis;

(o) providing for the taking of samples of any article for the purposes of this Act or for any other purpose;
(k) exempting any food, health from all or any of the provisions of this Act and prescribing the conditions of such exemption;

(p) for the method of clearance of the articles regulated by this Act from the ports;

(q) prescribing forms and particulars to be provided in forms;

(r) governing donation and disposal of drugs, medical devices, therapeutic cosmetics or scheduled substances; and

(s) governing generic substitution;

(t) prohibiting, regulating or restricting the sale of specified health product or technology by any of the persons licenced under this Act or by any class of those persons;

(u) exempting from any of the provisions of this Act relating to the sale of any article;

(v) prohibiting, regulating or restricting the manufacture, sale or advertising of health products or technologies;

(w) the safe custody and storage of health products or technologies;

(x) prescribing the procedure for the declaration of commercial interest by members of the Board and Committees of the Authority;

(y) governing the electronic sale of medicines

(z) providing for the categorization and classification of medical devices;

(aa) governing administration of clinical trials of drugs and any other health product and technology;

(bb) governing the compiling, processing, keeping, and submission of the information on donation, collection, testing, processing, distribution, transfusion and other use, exportation, importation and destruction of blood and blood products; the containers in which scheduled substance health products may be supplied;
(cc) the addition to scheduled substances of specified ingredients for the purpose of rendering them readily distinguishable as scheduled substances;

(dd) prohibiting the sale by retail of a specified medicine, medical device or Scheduled Substance except on prescription given by a qualified medical practitioner, dentist or veterinary surgeon and for prescribing the form and regulating the use of those prescriptions;

(ee) providing for the manner in which a pharmacist or a person otherwise authorized under this Act may dispense medicines or medical devices;

(ff) provide the manner and procedure in which clinical trials may be conducted in Kenya;

(gg) the compounding of medicines and the dispensing of medicines and medical devices;

(hh) the period for which books or registers required to be kept for the purposes of this Act are to be preserved;

(ii) the fees to be paid for anything to be done under this Act;

(jj) a particular procedure to be observed by the Authority;

(kk) the conduct of inquiries by the Authority under this Act and the attendance of witnesses and the production of evidence thereat;

(ll) prescribing powers or duties to be performed or exercised by an analyst, methods of analysis or examination of samples for the purposes of this Act, the form of any certificate or report to be furnished in connection with such analysis or examination, or the nature or arrangement of particulars to be reflected in such a certificate or report;

(mm) generally for giving effect to this Act.
(2) The authority shall adhere to the principle of public participation in making regulations.

114. (1) Upon the date of coming into operation of this Act the former Board shall be dissolved and—

(a) all assets and liabilities of the former Boards shall be transferred to and vest in the Authority without further assurance and the Authority shall have all powers necessary to take possession of, recover and deal with such assets and discharge such liabilities;

(b) every agreement, whether in writing or not, and every deed, bond or other instrument to which the former Boards was a party or which affected the former Boards, and whether or not of such a nature that the rights, liabilities and obligations thereunder could be assigned, shall have effect as if the Authority were a party thereto or affected thereby instead of the former Boards and as if for every reference (however worded and whether express or implied) therein to the former Boards there were substituted in respect of anything to be done on or after such date of coming into operation a reference to the Authority.

(c) any proceedings pending immediately before such date of coming into operation to which the former Boards was a party shall be continued as if the Authority was a party thereto in lieu of the former Boards;

(d) all officers of the former Boards shall become the officers of the Authority and, subject to the provisions of any rules made under this Act, shall continue in office for the period for which they were appointed or elected as officers of the former Boards.

(2) In this section “the former Boards” means the Pharmacy and Poisons Board and the Board of the National Quality Control Laboratory established under the Pharmacy and Poisons Act, and, the Public Health (Standards) Board established under the Food and Drugs Act.
115. (1) The enactments specified in the second column of the Sixth Schedule are repealed to the extent specified in the third column of that Schedule.

(2) The provisions of this Act shall be in addition to and not in derogation of the provisions of the Public Health Act or its successor.
FIRST SCHEDULE  
(S. 14)
PROVISIONS AS TO THE CONDUCT OF BUSINESS AND
AFFAIRS OF THE BOARD

Meetings of the Board.

1. The Board shall meet at least four times in each year.

Special meetings.

2. The Chairperson may at any time convene a special meeting of the Board and shall do so within fifteen days of a written requisition for the meeting signed by at least three members.

Chairperson to preside.

3. (1) The Chairperson shall preside at all meetings of the Board, at which he is present and in the case of his absence, the Vice Chairperson shall preside.

(2) At a meeting of the Board at which neither the members of the Board present shall elect one of their numbers to preside, and the person so elected shall have all the powers of the chairperson with respect to that meeting and the business transacted thereat.

Quorum.

4. The quorum for the conduct of the business of the Board shall be nine members.

Voting procedure.

5. The decisions of the Board shall be by a majority of votes, and the Chairperson of the meeting shall have an original and a casting vote.

Validity of proceedings.

6. The validity of any proceedings of the Board shall not be affected by any vacancy among the membership thereof, or by any defect in the appointment of a member thereof.

Minutes.

7. Minutes of the proceedings at meetings of the Board shall be kept in such a manner as the Council directs, and, on the written request of the Cabinet Secretary, shall be made available to him or any person nominated by him.

Committees of the Board.

8. The Board may establish such committees as may be necessary for the performance of the functions of the Board and may, subject to the provisions of this Act, delegate powers conferred on it to any such committee.

Power of the Board to regulate own procedure.

9. Subject to the provisions of this Schedule, the Board shall regulate its own procedure.
10. (1) If a member of the Board is directly or indirectly interested in any contract, proposed contract or other matter before the Board and is present at a meeting of the Board at which the contract, proposed contract or other matter is the subject of consideration, he shall, at the meeting and as soon as reasonably practicable after the commencement thereof, disclose the fact and shall not take part in the consideration or discussion of, or vote on, any questions with respect to the contract or other matter, or be counted in the quorum of the meeting during consideration of the matter.

(2) A disclosure of interest made under this paragraph shall be recorded in the minutes of the meeting at which it is made.

11. A member of the Board or of an Advisory Committee appointed under this Act shall declare in writing upon appointment and at any time thereafter as applicable his or her commercial interests related to the pharmaceutical, food or health care industry, which interests shall include, but shall not be limited to, any consultancy, paid or unpaid, any research grant from which the member directly or indirectly benefits, or any equity holding or any executive or non-executive directorship or any other payment or benefit in kind, and shall recuse himself or herself from any discussion or decision-making to which the said interests relate or may relate.
SECOND SCHEDULE (s. 9)
OATH/AFFIRMATION OF THE OFFICE OF CHAIRPERSON/A MEMBER/DIRECTOR

I, ................................................................. having been appointed (the chairperson/member/director of) the Kenya Food and Drugs Authority under the Kenya Food and Drugs Administration Act, 2012, do solemnly (swear/declare and affirm) that I will at all times obey, act and uphold the Constitution of Kenya and all other written laws of the Republic, that I faithfully and fully, and impartially and to the best of my ability, will discharge the trust and affirm the functions and exercise the powers devolving upon me by virtue of this appointment without favour, bias, affection, ill-will or prejudice. (SO HELP ME GOD).

Declared by the said ..........................................................

Before me this ................. day of ...............................

..........................................................
Chief Justice

THIRD SCHEDULE (s.8)
PROVISIONS RELATING TO MEMBERS OF THE BOARD

1. (1) This paragraph provides for the appointment of a member of the Board nominated by a body under Section 8(2) (i) and (k) of this Act.

(a) The nominating body shall submit the names of the nominees to the Cabinet Secretary.

(b) being satisfied that the nominating bodies have complied with the conditions under the Act, the Cabinet Secretary shall by notice in the gazette appoint the said members.

2. This paragraph provides for the appointment of a member of the Board under section 10(2)(a) of this Act—
(a) A selection panel shall submit the names of the nominees to the Cabinet Secretary.

(b) being satisfied that the selection panel has complied with the conditions under the Act, shall submit the names of the nominees to the National Assembly for approval.

(c) The National Assembly shall, within fourteen days after it first meets after receiving the names of the nominees—

(i) consider the nominees and either approve them or reject all or any one of them; and

(ii) notify the Cabinet Secretary as to its approval or rejection under subparagraph (a).

(d) If the National Assembly approves a nominee, the Cabinet Secretary shall, within fourteen days after receiving the notification of the National Assembly, forward the name of the nominees to the President and the President shall, within fourteen days after receiving the name, appoint the nominee as a member of the Board.

(e) If the National Assembly rejects any of the nominees submitted by the nominating body, the Cabinet Secretary shall, within fourteen days after receiving the notification of the National Assembly, request the nominating body to submit a new nominee to the Cabinet Secretary and subparagraph (3), (4) and (5) and this subparagraph apply with necessary modifications with respect to that new nominee.

(f) In nominating and approving persons to be members of the Board, the selection panel and the National Assembly shall have regard to—

(i) the honesty and integrity of the person and the person’s knowledge and experience; and

(ii) the importance of representing Kenya’s gender, regional and other diversities on the Board.

(iii) Absence of any commercial interest in the pharmaceutical, veterinary, food or healthcare industry
(iv) any other provisions in this Act and in the Constitution.

(g) Within seven days after any vacancy arises in the membership of the Board, the Cabinet Secretary shall request the nominating body or the Selection Panel as the case may be to submit nominees under subparagraph (2) and the nominating body or selection panel shall do so within twenty-one days after being requested to

3. The following shall apply with respect to the initial appointment to the Board following the commencement of this Act—

(a) each nominating body shall submit its initial nominees within twenty-one days after the commencement of this Act.

(b) the Cabinet Secretary shall wait until sufficient nominees are approved to form a quorum before submitting the names of the approved nominees under subparagraph (5);

(c) within fifteen days after sufficient numbers of the members of the Board are appointed to form a quorum, the Cabinet Secretary shall call a meeting of the Board for the purposes of nominating the Chairman and the Vice-chairman.

4. (1) A person may not serve more than two terms as a member of the Board.

(2) A member of the Board shall, unless his office becomes vacant as provided for under this Act, continue to hold office until he is appointed or replaced by another member appointed under the Act.

5. The President, on the recommendation of the Board, may terminate a person's appointment as a chairperson of the Board only if the person—

(a) is unable to perform the functions of his office by reason of mental, or physical infirmity;

(b) is adjudged bankrupt;

(c) is convicted of a crime carrying a jail term of not less
than six months; or

(d) is absent from three consecutive meetings of the Board without reasonable excuse.

6. (a) A member of the Board who has direct or indirect personal interest in a matter being considered or to be considered by the Board shall, as soon as reasonably practicable after the relevant facts concerning the matter have come to his knowledge, disclose the nature of his interest to the Board.

(b) A disclosure of interest in a matter shall be recorded in the minutes of the meeting of the Board and the members shall not be present while that matter is being dealt with by the Board and shall not take part in any deliberations or vote relating to the matter.

7. The Authority shall pay the members of the Board such allowances and expenses as are determined by the Cabinet Secretary in charge of finance in consultation with a Committee of the National Assembly designated by the National Assembly for that purpose.

8. (a) No action or proceeding for compensation or damages shall be brought against a member of the Board or any other person authorized by the Board, in respect of anything done or omitted to be done in good faith under this Act.

(b) This paragraph shall not relieve the Board of any liability in law.

FOURTH SCHEDULE (s. 21(2))

SCIENTIFIC ADVISORY COMMITTEES

1. (1) There shall be a committee of the Authority, to be known as the National Food Safety Committee, which shall have the responsibility within the Authority for advising the Cabinet Secretary on the formulation, implementation and monitoring of the food safety policy.

(2) The Committee shall consist of the following members—

(a) a representative of the Kenya Bureau of Standards;
(b) a representative of the Kenya Agricultural and Livestock Research Organization;
(c) a representative of the Kenya Plant Health Inspectorate Services;
(d) a representative of the Department of Public Health;
(e) a representative of the Weights and Measures Department;
(f) a representative of the Government Chemist's Department;
(g) a representative of the Department of Veterinary Services;
(h) a representative of the Kenya Dairy Board;
(i) a representative of the Horticultural Crops Development Authority;
(j) the Deputy Director General for Foods of the Authority, who will act as Secretary to the Committee; and
(k) two members appointed by the Director General from among the staff.

(3) Of the two members appointed under subparagraph (2) (k)—

(a) one shall be a person with executive responsibility in the Authority for food inspections and

(b) one shall be a person with responsibility within the Authority.

(4) The members of the National Food Safety Coordination Committee shall hold office for a term of three years and shall be eligible for re-appointment for one further term of three years.

2. (1) There shall be a Committee of the Authority, to be known as the Human Medicines Committee, which shall have the responsibility within the Authority for advising the Authority on regulatory policy relating to human medicinal products, giving advice in relation to the safety, quality and efficacy
of these, and promoting the collection and investigation of information relating to adverse reactions involving these products.

(2) The Committee shall consist of the following members:

a Chairperson appointed by the Cabinet Secretary knowledgeable in matters of human medicines regulation;
the Deputy Director General for medicines who shall be the secretary to the committee;
two members with relevant knowledge, experience and expertise appointed by the Director General from among the Authority staff;
four other members who have knowledge, experience and expertise in matters relating to human medicines, healthcare and public health appointed by the Cabinet Secretary; and
the Ministry of Health Director of Health Products and Technologies (or equivalent), or his or her designate, who shall be a non-voting member.

(3) Of the two members appointed under subparagraph (2) (d) —

(a) one shall be a person with executive responsibility in the Authority for medicines evaluation and assessment; and
(b) one shall be a person with responsibility within the Authority for inspections related to human medicines

(4) Each member appointed under subparagraph (2) (a), (b) and (d) shall hold office for a term of three years and shall be eligible to be appointed for one additional term.

(5) The Committee shall elect a Vice-chairperson from amongst its members of a different gender to the Chairperson who person shall hold this office for a term of three years and shall be eligible to be appointed for one additional term.

3. (1) There shall be a Committee of the Authority, to be known as the Veterinary Medicines Committee,
which shall have the responsibility within the Authority for advising the Authority on regulatory policy relating to veterinary medicinal products, giving advice in relation to the safety, quality and efficacy of these, and promoting the collection and investigation of information relating to adverse reactions involving these medicines.

(2) The Committee shall consist of the following members—

(a) a Chairperson appointed by the Cabinet Secretary, in consultation with the Cabinet Secretary responsible for veterinary services, and who is knowledgeable in matters of veterinary medicines regulation;

(b) The Director of Veterinary Services or his or her designate;

(c) the Deputy Director General for Medicines who shall be the secretary;

(d) two members with relevant knowledge, experience and expertise appointed by the Director General from among the staff;

(e) four other members who have knowledge, experience and expertise in matters relating to medicines regulation, veterinary services and veterinary public health appointed by the Cabinet Secretary; and

(f) the Director of Health Products and Technologies, or his representative, who shall be a non-voting member.

(3) Of the two members appointed under subparagraph (2) (d)—

(a) one shall be a person with executive responsibility in the Authority for veterinary medicines evaluation and assessment

(b) one shall be a person with responsibility within the Authority for veterinary medicines related inspections.

(4) Each member appointed under subparagraph (2) (a), (b) and (e) shall hold office for a term of three years and shall be eligible to be appointed for one additional term.
4. (1) There shall be a Committee of the Authority, to be known as the Medical Devices Committee, which shall have the responsibility within the Authority for advising the Authority on regulatory policy relating to medical devices, giving advice in relation to the safety, quality and efficacy of these, and promoting the collection and investigation of information relating to any issues involving their use.

(2) The Committee shall consist of the following members—

(a) a Chairperson appointed by the Cabinet Secretary, in consultation with the Cabinet Secretary responsible for health, and who is knowledgeable in matters of medical devices specifications and use;

(b) The Director of Medical Engineering Services or;

(c) the Deputy Director General for Medicines who shall be the secretary;

(d) two members with relevant knowledge, experience and expertise appointed by the Director General from among the staff;

(e) four other members who have knowledge, experience and expertise in matters relating to medical devices regulation, nursing services and public health services appointed by the Cabinet Secretary; and

(f) the Director of Health Products and Technologies, or his representative, who shall be a non-voting member.

(3) Of the two members appointed under subparagraph (2) (d)—

(a) one shall be a person with executive responsibility in the Authority for medical devices evaluation and assessment; and

(b) one shall be a person with responsibility within the Authority for medical devices related inspections.

5. (1) There shall be a Scientific Committee of the Authority, to be known as the National Quality
Control Committee, which shall have the responsibility for advising the Authority and Cabinet Secretary on quality control of health products and promoting the collection and investigation of information relating to quality of health products, maintaining relevant standards and certifications; and related matters necessary to ensure proper and efficient running of the National Quality Control Laboratory.

(2). The National Quality Control Committee shall consist of the following members—

(a) a chairman appointed by the Cabinet Secretary, and who is knowledgeable in matters of quality control

(b) the Deputy Director General for medicines or his or her designate;

(c) the Deputy Director General for foods or his or her designate;

(d) four other members who have knowledge, experience and expertise in matters relating to health products regulation, and quality control of medicines, therapeutic foods and therapeutic cosmetics, and medical devices; and

(e) the Director of the National Quality Control Laboratory, who shall be the secretary.

FIFTH SCHEDULE

The current editions of:

<table>
<thead>
<tr>
<th>Name</th>
<th>Abbreviation</th>
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<tbody>
<tr>
<td>Pharmacopoeia Internationalis .. ..</td>
<td>(Ph.I.)</td>
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<tr>
<td>The British Pharmacopoeia .. ..</td>
<td>(B.P.)</td>
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<tr>
<td>The Pharmacopeia of the United States of America</td>
<td>(U.S.P.)</td>
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<tr>
<td>Codex Francais .. .. ..</td>
<td>(Codex)</td>
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<tr>
<td>The Canadian Formulary .. ..</td>
<td>(C.F.)</td>
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<tr>
<td>The British Pharmaceutical Codex .. ..</td>
<td>(B.P.C.)</td>
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</tbody>
</table>
SIXTH SCHEDULE (s. 71)
PURPOSES FOR WHICH DRUGS, MAY NOT BE ADVERTISED

1. The cure of syphilis, gonorrhoea or soft chancre in any of their forms.

2. The prevention, relief or cure of Bright’s disease, schistosomiasis, cancer, Human Immunodeficiency Virus infection/Acquired Immune Deficiency Syndrome, consumption or tuberculosis, leprosy, lupus, diabetes, epilepsy or fits, locomotor ataxy, paralysis, or infantile paralysis.

3. The cure of arterio-sclerosis, septicaemia, diphtheria, dropsy, erysipelas, gallstones, kidney stones and bladder stones, goiter, heart disease, tetanus or lockjaw, pleurisy, pneumonia, scarlet-fever, smallpox, trachoma, amenorrhoea, hernia or rupture, blindness or any structural or organic ailment of the auditory system.

4. The cure of any habit associated with sexual indulgence, or of any ailment associated with those habits; or the restoration or stimulation of the sexual functions.

SEVENTH SCHEDULE (s.116)

<table>
<thead>
<tr>
<th>Number</th>
<th>Short title</th>
<th>Extent of Repeal</th>
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</thead>
<tbody>
<tr>
<td>Cap 244</td>
<td>The Pharmacy and Poisons Board Act</td>
<td>The whole Act</td>
</tr>
<tr>
<td>Cap. 254</td>
<td>Food, Drugs and scheduled substances Act</td>
<td>The whole Act</td>
</tr>
<tr>
<td>No. 14 of 1994</td>
<td>Narcotic Drugs and Psychotropic Substances Act</td>
<td>Sections 16, 17 and 18</td>
</tr>
</tbody>
</table>
MEMORANDUM OF OBJECTS AND REASONS

Statement of the Objects and Reasons for the Bill

The regulation of medicines, pharmaceutical practice, food, drugs, scheduled substances, therapeutic cosmetics, medical devices and related substances has been in fragmented legislation. In keeping with international best practices and guidance from the World Health Organization, the aim of the proposed national health products regulatory system is to safeguard the health of the public by ensuring the quality, safety and efficacy/effectiveness of medicines, food and related health products; based on principles of sound science, evidence and transparency. This requires an institutional framework with a clear legal mandate; and with the requisite expertise and independence in decision-making. It is for these main reasons that the Kenya Food and Drugs Regulation Bill is proposed. The proposed Authority will be established within National Government, its functions being a critical and integral part of Health Policy, as set out in the Fourth Schedule to the Constitution.

Clause 2 of the Bill provides for the definition and interpretation of terms used in the Bill.

Clause 3 of the Bill further provides where the Act shall apply regarding the regulation of health products and technologies including:

(a) Blood and blood products
(b) Chemical substances
(c) Therapeutic cosmetics
(d) Food
(e) Herbal medicines and products
(f) Medical devices including radiation-emitting devices;
(g) Medicines;
(h) Scheduled substances;
(i) Tobacco and tobacco products

Clause 4 of the Bill establishes the Kenya Food and Drugs Authority as a corporate body capable of suing and being sued; taking, purchasing, acquiring, holding or disposing of movable or immovable property; and doing or performing all such other things or acts for the proper discharge of its functions under the Act, which may be lawfully done by a body corporate.

Clause 6–7 of the Bill establishes the office of the Director General of the Authority. The Clause further establishes the qualifications for
appointment us Director General of the Authority as well as the function of the Director General.

Clause 8 of the Bill establishes the management of the Authority. It stipulates who shall sit on the Board of the Authority. It further states who shall qualify to be on the Board of the management.

Clause 9 of the Bill provides for the oath of office.

Clause 10 of the Bill states when the office of the chairperson or member of the Board shall become vacant.

Clause 11 of the Bill states the procedure for removing a member of the Board and such application shall be copied to the Chairperson.

Clause 12–14 of the Bill provide for the functions and the powers and affairs of the Authority.

Clause 21 of the Bill provides for the establishment and functions of scientific advisory committees. These committees shall be established as necessary for the performance of the cabinet secretary’s functions and powers under this Bill.

Clause 22 of the Bill stipulates that the Authority shall regulate and monitor the manufacture, processing, distribution, sale and importation of food in Kenya.

Clause 23 of the Bill provides for the various types of foods that shall not be sold in Kenya. These include foods that: contain poisonous or harmful substances, unfit for human consumption, contains rotten or diseased substances or is adulterated.

Clause 24 of the Bill provides that a person who has reason to believe that any food that is not in compliance with the provisions of this Act has a duty to notify the Authority.

Clause 25-26 of the Bill provide that any person who advertises, labels or sells any food or labels, packages any food in contravention with this Act commits an offence.

Clause 27 of the Bill declares the act of selling any food that is below the nature, substance or quality demanded commits an offence

Clause 28 of the Bill provides for the foods that shall not be imported into Kenya including unsafe food and food that has been treated with or that contains a prohibited substance.

Clause 31 of the Bill provides for when and how a regulatory officer may examine food intended for human consumption which has been sold or is exposed for sale. This Clause further provides for the actions that shall be
taken by the regulatory officer in the event that the food is unfit for human consumption.

Clause 32–33 of the Bill concern Therapeutic foods. These Clauses classify therapeutic food as well as establish a Therapeutic Food Register.

Clause 34 of the Bill provides the prohibition of a number of medicines. This Clause further declares the offence and punishment for any person who sells any medicine that is prohibited under this Act or any other written Law. It also provides for an exception to this rule in the case of medicine compounded by a pharmacist or pharmaceutical technologist.

Clause 36 of the Bill provides for various offences that can be committed in respect to the standard of medicines including their content, packaging and advertising.

Clause 37-38 of the Bill ensure that the sale and preparation of medicines is up to standard by declaring it an offence to sell medicine that is not of the correct standard or preparing medicine in a manner that does not meet prescribed standards.

Clause 39 of the Bill detail the factors that the Authority shall take into consideration before granting a product licence.

Clause 40 of the Bill establishes a Medicines Register.

Clause 41-45 of the Bill provides for the process of registration of medicines and medical devices, these clauses also detail the process of amendment, transfer and cancellation of registration.

Clause 46 of the Bill provides for measures that the Cabinet Secretary shall be able to pursue in order to ensure supply of more affordable medicines.

Clause 47 of the Bill provides for when and how a pharmacist can dispense interchangeable multi-source medicine instead of what was prescribed by a medical doctor or dental practitioner.

Clause 48 of the Bill regulates the sale, manufacture, exportation and importation of herbal medicines.

Clauses 59-60 of the Bill provide for the formalities relating to licences for the manufacture of medicinal substances.

Clause 61 of the Bill provides for the types of therapeutic cosmetics whose sale is prohibited and makes the sale of any such therapeutic cosmetics an offence.

Clause 62 of the Bill prohibits the sale of therapeutic cosmetics in such a manner as to make it likely for a therapeutic cosmetic that is not up to set
standards to be confused for another therapeutic cosmetic that is up to the
standards and makes it an offence to do so.

Clause 63 prohibits the sale of therapeutic cosmetics under insanitary
conditions and makes it an offence to do so.

Clause 64 provides for the treatment of any therapeutic cosmetic which
either contains a Scheduled Substance or claims to have a therapeutic
effect as a medicine.

Clause 65 of the Bill provides for the creation of a therapeutic cosmetics
register.

Clause 66 of the Bill provides for the prohibition of ingredients contained
in therapeutic cosmetics by the Authority and makes the sale of any
cosmetics containing these prohibited ingredients an offence.

Clause 67 of the Bill provides for the creation of a register containing all
duly registered medical devices and which shall include all the particulars
of the devices and the holders of the devices, that this Bill or any other law
requires to be contained in it.

Clause 68 of the Bill provides for the types of medical devices whose sale
is prohibited and makes the sale of any such medical devices an offence.

Clause 69 of the Bill prohibits the sale of any medical device in a manner
that is deceptive regarding any of the attributes of the medical device and
makes it an offence to do so.

Clause 70 of the Bill provides for the issuance of standards by the
Authority to ensure that the exposure of patients to radiation is minimised
and prohibits the sale of any medical device that does not conform to these
standards in such a manner that it is likely to be confused for a medical
device that does conform to the standards.

Clause 71

of the Bill provides for certain offences relating to the sale of medical
devices, including:

(a) Sale of any medical device under insanitary conditions.

(b) Sale of any medical device that has a measuring function that
does not provide accurate measurements.

(c) Sale of unapproved medical devices

Clause 72 of the Bill provides for the licensing of persons that deal in
blood and blood products setting out the manner in which a licence may be
obtained, registered and revoked.
Clause 73 of the Bill provides for the collection of blood in exchange for payment and establishes a register for blood and blood products containing healthcare establishments that deal with blood and blood products and serious incidents associated with the collection and use of blood and blood components.

Clause 74 of the Bill provides for the manner in which the Authority shall handle serious and adverse incidents relating to blood and blood components.

Clauses 75–77 of the Bill provide for the manner in which the Authority shall regulate the manufacture of tobacco.

Clauses 78–79 of the Bill provide for the establishment of the National Quality Control Laboratory whose functions include:

(a) Examination and testing of drugs to ensure quality control drugs and medicinal substances

(b) Performing pharmaceutical evaluation

(c) Conducting research and training and testing the quality of locally manufactured and imported medicines, medical devices or therapeutic foods on behalf of the Authority with a view to determining whether they comply with set standards.

Clauses 80–87 of the Bill provide for the manner in which the Authority shall regulate the advertisement and labelling of health products and technologies.

Clause 88 of the Bill provides the Cabinet Secretary with the authority to prohibit or control certain medicines and makes the contravention of any order of prohibition made by the Cabinet Secretary in exercise of this authority an offence.

Clause 89 of the Bill provides for the situations under which the Authority may permit persons to supply health products or technologies that are not registered.

Clause 90 of the Bill provides for the retention and disposal of any drugs seized under this Bill.

Clause 91 of the Bill provides for the situations under which the Authority may compel a person handling health products to provide the Authority with information regarding the health products.

Clauses 92–95 of the Bill provide for the inspection of licences and certificates of registration by regulatory officers and makes the obstruction of regulatory officers from exercising their powers an offence.
Clause 96 of the Bill provides for the criminal responsibility of directors, secretaries and managers where an act that is an offence under this Act is committed by a body corporate and applies the same to partners.

Clause 97 of the Bill provides for the inspection of animals and seizure of meat meant for human consumption that is deemed to be unfit for consumption by regulatory officers.

Clause 98 of the Bill provides for the powers of regulatory officers and prescribes the manner in which they shall carry out their duties.

Clause 99 of the Bill provides for the powers of the Director of Medical Services, Director of Veterinary Services or the Director of Agriculture to order a public officer to have articles of food or health products analysed.

Clause 101 of the Bill provides for the power of the Cabinet Secretary to obtain the particulars of any substance.

Clause 102 of the Bill provides for the power of the court to order the cancellation of a licence issued under the Act.

Clause 103 of the Bill provides for the prosecution of any person for an offence under this Act.

Clause 104 of the Bill provides for the penalties to which a person shall be liable for offences committed under this Act.

Clauses 106-112 of the Bill contain financial provisions relating to the Authority including:

(a) The funds of the Authority
(b) The financial year of the Authority
(c) The preparation of the annual estimates of the Authority
(d) The keeping of accounts and auditing of the Authority
(e) The manner in which the Authority may invest its funds
(f) The preparation of annual reports by the Authority
(g) The submission of special reports to the National Assembly

Clauses 112-115 of the Bill contain miscellaneous provisions relating to the functioning of the Authority including:

(a) Making of regulations for the effective enforcement of the Act
(b) Transitional provisions
(c) The laws or sections thereof repealed by the enactment of this Act
Statement on the delegation of legislative powers and limitation of fundamental rights and freedoms

The Bill does not contain any provisions limiting any fundamental rights or freedoms.

Statement of how the Bill concerns county governments

The Bill affects the functions of County governments as set out in the Fourth Schedule to the Constitution and is therefore a Bill concerning counties.

Statement as to whether the Bill is a money Bill within the meaning of Article 114 of the Constitution

The enactment of this Bill shall occasion additional expenditure of public funds.

Dated the 5th March, 2019.

ROBERT PUKOSE,
Member of Parliament.