Development of an analytical tool to assess Biosecurity legislation

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Globalization of trade in agricultural products brings opportunities and risks. On the one hand, it generates wealth in countries exporting their produce to foreign markets and brings that produce to the tables of consumers in far-away lands. On the other hand, it opens new pathways for pests and diseases that can damage natural resources with accompanying economic and environmental consequences. In order to capture those opportunities and manage those risks, there is an increasing recognition of the need to integrate and improve coordination of regulatory activities designed to protect human, animal and plant life and health and the environment.

Interest in Biosecurity comes in response to these needs. It attempts to draw together the policy and regulatory frameworks for risk assessment and risk management across the sectors of food safety, animal life and health (including fisheries) and plant life and health. Biosecurity aims to manage biological risks in these three sectors while protecting the environment and contributing to its sustainable use. In essence, Biosecurity balances enthusiasm for international trade with the need to protect against risks. Transparent and efficient controls in these sectors need not create unnecessary barriers to international trade; rather they facilitate it.

Biosecurity is an interdisciplinary activity that covers a wide range of subjects and approaches. As Biosecurity works towards the integration of animal health, plant health and food safety in order to streamline risk assessment and risk management practices, the division of responsibilities among national agricultural regulatory authorities comes under scrutiny. Controls and authorities for Biosecurity matters tend to be scattered over a variety of ministries, including the ministries of agriculture, health, environment and trade and industry. The objective of Biosecurity is to draw together relevant regulatory authorities or to create coordinating mechanisms to streamline approaches to managing biological risks. To implement the necessary coordination, countries must look closely at their national legal frameworks. This will aid in implementing the most efficient institutional set-up while also protecting rights and establishing responsibilities in a way that is conducive to the active participation of public authorities, the private sector and consumers.
To implement a *Biosecurity* approach, governments should first identify and analyse the existing constellation of legal provisions covering the subject areas of *Biosecurity*. At times this may not be easy as *Biosecurity* is often regulated in a plethora of parliamentary-level and subsidiary pieces of legislation of different natures, scopes and objects. The present study elaborates an orderly methodology to facilitate the review and assessment of national legal frameworks for *Biosecurity*. The methodology arises from work carried out in six pilot countries – Ethiopia, Ghana, India, Kenya, Uganda and Viet Nam – by national legal experts actively testing and refining the analytical tool.

The methodology set out in this study should enable a comprehensive evaluation of national laws and regulations covering the main subject areas of *Biosecurity*, comparing national rules with international requirements and providing an overall evaluation of the national regulatory framework *vis-à-vis* the objectives of *Biosecurity*. It is hoped that the methodology will be a useful tool for countries wishing to assess and develop updated legislation to achieve a *Biosecurity* approach.

A number of people have participated in the development of this legal analytical tool. Daniele Manzella and Jessica Vapnek were the principal authors. International legal consultants Emmanuelle Bourgois and Charlotta Jull provided discrete inputs, while Ariella D’Andrea, Jennifer Hilton, Abdul Rahman Lediju, Victor Mosoti, Valerio Poscia and George Sarpong provided excellent research assistance or comments on the overall methodology. Wondwossen Sintayehu Wondemagegnehu (Ethiopia), George Sarpong (Ghana), Roopa Madhav and Adil Hasan Khan (India), Patricia Kamerni-Mbote (Kenya), Judy Obitre-Gama (Uganda) and Duong Thanh An (Viet Nam) carried out the national legal studies which formed the basis of the methodology and which are set out in Chapters 5–10. Niek Van der Graaff and Mike Robson provided key support for the concept and enabled the effective execution of the project. Essential funding was provided by the Government of Norway under the Programme Cooperation Agreement for *Biosecurity*.
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INTRODUCTION

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I. OVERVIEW

In 2006, the Government of Norway and the Food and Agriculture Organization of the United Nations (FAO) entered into a Programme Cooperation Agreement (PCA) to work on food security and poverty reduction, policy assistance and capacity building in low-income developing countries.

The fundamental goal of the FAO-Norway programme of cooperation is to alleviate hunger. In pursuance of this goal, the PCA seeks to help developing countries improve national capacities to meet domestic and international marketing requirements, decrease biological risks and improve preparedness for food crises that increase the risk of market collapse. The PCA activities address several of the UN Millennium Development Goals, aiming to eradicate hunger and poverty, ensure environmental sustainability and develop an open trading system that is rule-based, predictable and non-discriminatory.

Biosecurity\(^\text{1}\) draws together the policy and regulatory frameworks for risk management across the sectors of food safety, animal life and health (including fisheries) and plant life and health. The approach aims to manage biological risks in these three sectors while protecting the environment and contributing to its sustainable use.

Within the area of Biosecurity, the PCA programme has activities in:

- animal health;
- food safety;
- plant health;
- fish product safety;
- socio-economic analysis;
- policy development; and
- development law.

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\(^{1}\) Because translation of the word "biosecurity" into French and Spanish can lead to confusion, FAO capitalizes and italicizes it when referring to this regulatory approach in these three official FAO languages. See Biosecurity in Food and Agriculture, FAO Committee on Agriculture, 17th Session, 31 March–4 April 2003, Rome.
An important result of improving Biosecurity is that it enables countries to participate in an increasingly standards-driven international food and agricultural trading market, which is one of the key means of alleviating poverty in developing countries.

Biosecurity is an interdisciplinary subject, and thus the projects and studies under the umbrella of the PCA cover a wide range of disciplines and approaches. Along with this legal study, other projects undertaken through the PCA include testing of the Biosecurity Capacity Assessment Tool;\(^2\) defining data items for animal health/Biosecurity country profiles; improving support for FAO’s crisis management; carrying out studies in East Africa on district-level Biosecurity problems; and preparing studies on market collapse, fish product safety, aquatic animal health and the socio-economic aspects of Biosecurity. All activities take into account social, economic and gender issues; focus on poverty alleviation; and are undertaken in strategic cooperation with relevant regional groups, international organizations and other partners.

Under the PCA, the Legal Office proposed to develop an analytical tool to assess national Biosecurity legal frameworks. The tool consists of a methodological examination of the national laws and regulations covering the main subject areas of Biosecurity, comparing national rules with international requirements and providing an overall evaluation of national regulatory and institutional frameworks vis-à-vis the objectives of Biosecurity. The analytical tool is designed either for stand-alone use or for use with the Biosecurity Capacity Assessment Tool.

The programme of work for the development of this analytical tool consisted of the following activities:

1. identifying six low-income countries in different regions as the pilot countries for the activity;
2. recruiting one national legal consultant for each country to analyse the existing legislation on Biosecurity and the institutional structures for its implementation;

\(^2\) The Biosecurity Capacity Assessment Tool (currently in draft form) assists in assessing Biosecurity capacity needs across all sectors and all sector organizations at national level. It is a part of a larger Biosecurity tool kit and focuses on policy, legal and institutional frameworks, infrastructure and operations, risk analysis, technical capability and information exchange.
(3) drawing together the six analyses and developing the methodology for use in subsequent country assessments;
(4) editing and preparing the results for publication.

The first two countries selected as pilot case studies to develop the methodology were Ghana and Kenya, and additional case studies in Ethiopia, India, Uganda and Viet Nam followed. The lead author carried out national consultations in Ghana, Kenya and Uganda, with the support of the respective national legal consultants. The purpose of the consultations was to discuss the findings of the national legal consultants’ reports and the feasibility of legislative change.3

II. DEFINITION OF BIOSECURITY

As noted above, Biosecurity is a strategic and integrated approach that encompasses the policy and regulatory frameworks to analyse and manage risks in the sectors of:

- food safety;
- animal life and health; and
- plant life and health, including associated environmental risks.

These sectors include:

- food production in relation to food safety;
- the introduction of plant pests, animal pests and diseases and zoonoses;
- the introduction and release of genetically modified organisms and their products; and
- the introduction and safe management of invasive alien species and genotypes.

The objective of Biosecurity is to identify, assess and respond appropriately to all pests and diseases posing a significant threat to agriculture, forestry,

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3 The national consultations took place in Ghana from 22 to 26 January 2007, in Kenya from 30 April to 4 May 2007 and in Uganda from 5 to 11 May 2007.
horticulture, fisheries, native biodiversity and human health. Appropriate responses include eradication, containment and on-going control.

III. INTERNATIONAL CONTEXT

Biosecurity is of growing interest as a result of developments at the international level, including globalization of the world economy, technological progress and the rapid increase in communications, transport and trade. Against this background, there is concern that the appropriate level of protection of human, animal and plant life and health is not being maintained as risks increase.

The term Biosecurity does not appear in any instrument of international law. But as will be described in greater detail in the next chapter, the main international regulatory instruments and organizations that led FAO to adopt the concept and promote a specific work programme in relation to the Biosecurity approach are:

- the World Trade Organization Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement);
- the Convention on Biological Diversity (CBD) and the Cartagena Protocol on Biosafety (Cartagena Protocol);
- the Codex Alimentarius Commission (Codex);
- the Office international des épizooties (OIE, or World Organization for Animal Health); and
- the International Plant Protection Convention (IPPC).

The SPS Agreement identifies the rights of states concerning sanitary and phytosanitary measures in relation to international trade and also provides common obligations that govern those rights. Sanitary measures are those designed to protect animal and human life and health, while phytosanitary measures refer to the life and health of plants. The SPS Agreement provides for a unified approach to the different sectors of Biosecurity. The approach is

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4 For an analysis of the major international instruments relevant to Biosecurity, see A. Ingrassia, International and Regional Regulatory Frameworks Relevant to Biosecurity for Food and Agriculture, paper presented at the FAO Technical Consultation on Biological Risk Management in Food and Agriculture (unpublished).
centred on harmonization through international standards, science-based risk assessment and minimization of interference with international trade.

While traditional sanitary and phytosanitary controls were designed to ensure efficient production through the protection of natural resources, modern controls tend to integrate these concerns into a wider spectrum of issues, such as preservation of the environment and protection against the loss of biodiversity. Increasing awareness of these threats has expanded the scope of Biosecurity from its traditional focus on protection of primary production and trade. Under the SPS Agreement, three organizations – Codex, the IPPC and the OIE – are recognized as the sources of international standards for food safety, plant life and health and animal life and health, respectively.

The SPS Agreement recognizes the right of countries to take emergency measures based on incomplete information. In that respect, the agreement is complemented by the Cartagena Protocol,\(^5\) which is based on the precautionary principle. In this context the principle provides that, where an activity increases the threat of harm to human health or the environment, precautionary measures should be taken even if some causal relationships are not fully established scientifically.

Other international instruments can be said to form part of the Biosecurity-related regulatory framework. For instance, the OIE Terrestrial Animal Health Code, the Aquatic Animal Health Code and their respective Manuals for Diagnostic Tests outline import and export procedures to avoid disease spread and structures for the communication of epidemiological information. Several Codex documents are also relevant, including the Principles for Food Import and Export Certification and Inspection; Guidelines for the Design, Operation, Assessment and Accreditation of Food Import and Export Inspection and Certification Systems; and Guidelines for the Exchange of Information between Countries on Rejections of Imported Food.

In addition, some IPPC standards (more precisely, some International Standards on Phytosanitary Measures – ISPMs) elaborate on environmental

\(^5\) The objective of the Cartagena Protocol is to ensure an adequate level of protection in the field of the safe transfer, handling and use of living modified organisms possessing a novel combination of genetic material obtained through the use of modern biotechnology (art. 1).\)
considerations and are relevant to the management of invasive alien species under the Convention on Biological Diversity (CBD). Article 8(h) of the CBD requires contracting parties to prevent the introduction of, and control or eradicate, those alien species which threaten ecosystems, habitats or species.\(^6\)

The multiple impacts of invasive alien species (IAS) call for coordinated international action to minimize their environmental as well as economic effects. Toward this end, the CBD and IPPC have been working cooperatively in several ways. The CBD Conference of the Parties and the IPPC have collaborated on the preparation of a supplement to ISPM No. 11 (Pest risk analysis for quarantine pests, including analysis of environmental risks and living modified organisms) in order to incorporate risks to biodiversity posed by IAS that are considered plant pests. Further collaboration has taken place in the revision of ISPM No. 3 (Guidelines for the export, shipment, import and release of biological control agents and other beneficial organisms) in order to manage risks to biodiversity that beneficial organisms may generate.

ISPM No. 11 includes the analysis of risks that living modified organisms (LMOs) present. In this regard, the IPPC standard is relevant to the regime regulating LMOs under the Cartagena Protocol. The protocol establishes an informed agreement procedure for ensuring that countries are provided with information in advance, including an assessment of risks to biological diversity, necessary to make informed decisions before agreeing to the import of such organisms into their territory. In the assessment of risks to biological diversity, ISPM No. 11 can be applied for LMOs that are categorized as plant pests.

Risk analysis is the basis for the establishment of sanitary and phytosanitary measures for the import of plants, animals and foods, and the concepts are the same across these sectors. Thus, risk analysis is one common thread among the many international instruments relevant to Biosecurity. But although international standard-setting and cooperation are important, the establishment, implementation and monitoring of Biosecurity in agriculture is a matter for national governments. How to implement a Biosecurity approach at national level is the subject of the next section.

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\(^6\) Under the CBD, an alien species is defined as "a species, subspecies or lower taxon, introduced outside its natural past or present distribution" and an invasive species is "an alien species whose introduction and/or spread threaten biological diversity" (art. 8(g)).
IV. IMPLEMENTATION OF BIOSECURITY AT NATIONAL LEVEL

The ultimate objectives of Biosecurity at the national level are to protect domestic agricultural production and natural resources from biological hazards and to safeguard the health of consumers in the food chain. To comply with the SPS Agreement, risk assessment in accordance with applicable international standards or scientific justification shall underlie domestic decision-making regarding the import and use of plants, animals and foods.

Countries require strong global and regional relationships to identify and manage emerging risks and this international network in turn supports appropriate national actions such as:

- comprehensive, competent surveillance programmes and diagnostic services to detect and identify the arrival and spread of pests and diseases;
- sufficient capability to conduct timely assessments of threats from new species;
- rapid response capability to eradicate new pests and diseases before they establish and spread; and
- standardization of science-based identification of all risk pathways and high-risk organisms, and implementation of pre-border and border measures to prevent pests and diseases from entering the country.

In order to enforce effective controls and to comply with international standards, countries need to build capacity in their administrations. There are several components of national capacity building which may assist countries in reducing unjustified obstacles to trade while protecting food safety, animal and plant life and health. These include developing national infrastructure, enhancing specific expertise and strengthening personnel and training.

In some countries (e.g. Belize and New Zealand), capacity building has concentrated on institutional aspects with a view to achieving the integration of the animal health, plant health and food safety sectors. The objective is to draw together relevant authorities and ministries in charge of these three sectors, or at least to create coordinating mechanisms. This tendency derives from the fact that responsibility for Biosecurity matters tends to be scattered over a variety of ministries, including the ministries of agriculture, health, environment and trade.
Coordination among the relevant authorities and ministries will improve outcomes with respect to activities such as:

- participation in the meetings of international standard-setting bodies for the definition of common international standards;
- exchange of relevant official information;
- allocation of national resources and capacities;
- input of scientific advice into all levels of policy, planning and decision-making;
- technical support of stakeholders across the spectrum of Biosecurity interests;
- elaboration of effective education and awareness programmes to encourage compliance with legislation; and
- enforcement of legislation.

Before coordination can take place, however, there is a need to assess existing policies and legislation and the allocation of responsibilities among the different institutions involved with agricultural trade. Government policies determine the desired levels of Biosecurity protection while laws and regulations outline how that protection will be achieved. Good policies and laws can create an environment conducive to the application of Biosecurity by the government and the private sector, including farmers and other small-scale producers and the commercial agro-food supply chain.7

V. LEGISLATIVE REVIEW AND ASSESSMENT

The general objectives of legislation are to protect rights and establish responsibilities as well as to enable the meaningful participation of all stakeholders, from central institutions to local communities. Good legislation establishes predictable rules for the exercise of public powers, which can encourage investment and facilitate the operation of markets while protecting public interests such as the conservation of natural resources.

Before a government can develop new legislation or amend the legislation in place, however, it must identify and analyse the existing constellation of legal

provisions covering the relevant subject areas. In other words, it is essential to know what the legislation says and to understand how the system operates under that legislation before making recommendations for change. An assessment of national legislation on *Biosecurity* should evaluate both compliance with international obligations and the allocation of roles and responsibilities of sectoral bodies in the management of biological risks for food and agriculture.

In some cases, where there are no laws or regulations on some or all of the elements of *Biosecurity*, entirely new legislation must be drafted. In other cases, there may be an existing legal framework but it may be outdated or insufficient, or rife with overlaps and gaps, and thus call out for a complete overhaul. In still other cases, only minor changes may be necessary, for example to add a few specific obligations or to enhance coordination among government bodies.

Effective institutional coordination avoids duplication, inconsistency and disputes among the relevant agencies and also helps improve efficiency in the application of sanitary and phytosanitary measures. The ultimate goal of upgrading national legal frameworks to regulate, manage and control *Biosecurity* for food and agriculture is to implement effective controls, increase cost effectiveness and improve consistency across sectors. Of course, if the analysis of the existing framework leads to the determination that the current legislation is good enough, time is better spent on other matters such as improving implementation and enforcement of existing laws.8

The next chapter provides an overview of the international framework for *Biosecurity*, while Chapter 3 presents the results of the gap analysis of the legislative frameworks for *Biosecurity* in the countries reviewed under the auspices of this project. Chapter 4 proffers a suggested analytical methodology to assess national *Biosecurity* legal frameworks and design an appropriate legal strategy for their improvement. Chapters 5 to 10 contain the national case studies while Chapter 11 offers some concluding observations.

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I. INTRODUCTION

As described in Chapter 1, Biosecurity, according to the FAO official definition, comprises three sectors: food safety, plant life and health and animal life and health. These sectors include the introduction and release of genetically modified organisms (GMOs) and the introduction and safe management of invasive alien species (IAS). The international legal framework for Biosecurity encompasses all international instruments governing these sectors, as well as instruments relevant more generally to management of risks associated with food and agriculture. The embrace of a Biosecurity approach at national level calls for the harmonization of national legislation with these international instruments.

This chapter examines the relevant international instruments to understand their content and the main obligations they generate in the main Biosecurity sectors. Considering that these sectors intersect with and, to some extent, are shaped by the international trade regime of the World Trade Organization (WTO), the chapter starts by presenting that overarching regime. The analysis of the international regulatory framework for food safety, animal health, plant protection, GMOs and IAS follows. The chapter concludes with a brief overview of other international instruments that are also relevant to Biosecurity, in that they address the management of risks associated with food and agriculture.

II. WTO AGREEMENTS

2.1. SPS Agreement

The WTO Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement)\(^1\) aims to prevent the use of sanitary and phytosanitary measures (SPMs) as disguised barriers to international trade and is binding upon all WTO member states. According to Annex A of the agreement, SPMs are defined as any measures applied to:

• protect animal or plant life or health from risks arising from the entry, establishment or spread of pests, diseases or disease-carrying organisms;
• protect human or animal life or health from risks arising from additives, contaminants, toxins or disease-causing organisms in foods, beverages or feedstuffs;
• protect from risks arising from diseases carried by animals, plants or products thereof, or from the entry, establishment or spread of pests; or
• prevent or limit other damage from the entry, establishment or spread of pests.

The SPS Agreement is the cornerstone of *Biosecurity*, attempting to strike a balance between the protection of human, animal and plant life and health on the one hand and the removal of barriers to international trade on the other. It establishes that SPMs may be applied only to the extent necessary to protect human, animal and plant life or health and must be based on scientific principles and sufficient scientific evidence (art. 2.2). Countries are obligated to ensure that their SPMs do not arbitrarily or unjustifiably discriminate between countries where identical or similar conditions prevail, and more fundamentally, SPMs shall not be applied in a manner which would constitute a disguised restriction on international trade (art. 2.3).

The SPS Agreement allows countries to set their own level of protection based on the assessment of risks to human, animal and plant life and health, and to establish SPMs in accordance with that level of protection (art. 5.1). However, the agreement encourages countries to apply international standards where they exist, and in Annex A identifies the official international standard-setting bodies. As long as a WTO member state employs international standards in the formulation of its national SPMs, these are presumed to be consistent with the provisions of the SPS Agreement (arts. 3.1 and 3.2).

Nonetheless, countries may adopt measures which result in a higher level of protection than that offered by an international standard, guideline or recommendation. In such cases, a WTO member state may be asked to provide scientific justification or to demonstrate that it had to depart from the relevant international standard because applying it would not have resulted in the level of protection the country considered appropriate (art. 3.3).
The SPS Agreement is designed to improve the transparency of SPMs, by requiring WTO member states to notify other countries of any new or changed sanitary and phytosanitary requirements which affect trade. The SPS Agreement requires each member state to establish an office (a so-called SPS Enquiry Point) to provide advance notice of any new or changed SPMs, thus giving other member states an opportunity to comment on them and facilitating information-sharing.2

In the event that available scientific evidence to justify a measure is insufficient, the SPS Agreement provides some flexibility for member states to adopt SPMs provisionally. Provisional SPMs can be adopted on the basis of "available pertinent information" derived from a variety of sources. However, member states must subsequently seek additional information to objectively assess the risk further and to review the SPM within a reasonable period of time (art. 5.7).

For the first phase of implementation (until the year 2000), developing and least developed countries, which make up about two-thirds of the WTO membership, were accorded special and differential treatment under the SPS Agreement (art. 10).3

2.2. TBT Agreement

The WTO Agreement on Technical Barriers to Trade (TBT)4 is an instrument that is peripheral to the SPS Agreement, which seeks to ensure that technical regulations and standards do not create unnecessary obstacles to international trade. Such technical regulations and standards include packaging, marking and labelling requirements.

The TBT Agreement does not apply to sanitary and phytosanitary measures as defined in Annex A of the SPS Agreement (art. 1.5). Rather, it applies to unsafe products which may have an effect on human, plant or animal life and health based on their packaging, marking and labelling. In that regard, it is relevant to Biosecurity.

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2 See article 7 and Annex B.
Under the TBT Agreement, WTO member states are required to use international standards whenever they impose technical regulations on products that are covered by the agreement (art. 2.4). Whenever a technical regulation is based on an international standard and is prepared, adopted or applied with respect to one of the legitimate objectives listed in article 2.2, it is "rebuttably presumed not to create an unnecessary barrier to trade" (art. 2.5). Nonetheless, member states can deviate from these international standards so long as they still fulfil one or more of the enumerated legitimate objectives under article 2.2. Technical regulations cannot be more trade-restrictive than necessary to fulfil the legitimate objectives (art. 2.2).

In the event that an international standard does not exist, or the technical content of a proposed technical regulation is not in accordance with the technical content of an existing international standard, and the technical regulation may have a significant effect on trade, the TBT Agreement requires the member state to engage in consultations with other member states (art. 2.9). The required steps are set out in the TBT Agreement (e.g., written justification; notice; notification through the secretariat; making copies available; reasonable time for comments) (arts. 2.5 and 2.9). However, some of these steps may be omitted in emergency situations (art. 2.10).

Annexed to the TBT Agreement is a Code of Good Practice for the Preparation, Adoption and Application of Standards to guide the development of standards in WTO member states. Standardizing bodies must not act contrary to or inconsistently with the code (art. 4.1). The application of standards by member states is premised on the same principles of international trade as technical regulations: national treatment (i.e., treatment of products originating in the territory of any other WTO member in a manner no less favourable than that accorded to like products of national origin); non-discrimination (i.e., equal treatment to products originating in the territory of any other WTO member no less favourable than that accorded to like products originating in any other member country); proportionality (i.e., measures should be no more strict than necessary) and avoiding unnecessary obstacles to trade (art. 5.1.2). Likewise, there is a preference for deriving national standards from international standards, guidelines and recommendations (art. 5.4).

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5 "Legitimate objectives" include national security requirements, the prevention of deceptive practices and the protection of human health or safety, animal or plant life or health or the environment.
III. PLANT HEALTH

The main international instrument regulating plant health is the International Plant Protection Convention (IPPC). The IPPC was adopted in 1951 and revised twice, in 1979 and in 1997. The 1997 New Revised Text came into force in October 2005 and is binding upon all contracting parties. The IPPC is a multilateral treaty whose main purpose is to secure "common and effective action to prevent the spread and introduction of pests of plants and plant products and to promote appropriate measures for their control" (art. I.1). "Pest" is broadly defined in the convention as "any species, strain or biotype, animal life or any pathogenic agent injurious or potentially injurious to plants or plant products" (art. II.1). The IPPC’s scope is broad enough to include the potential impacts of plant pests on the environment and the importation of living modified organisms that may directly or indirectly affect plants or other organisms. There is therefore potential for overlap with the Convention on Biological Diversity (CBD) and its Cartagena Protocol, which has led to growing cooperation between the two agreements.

The IPPC identifies modern phytosanitary concepts, such as pest risk analysis and the designation of pest free areas, and embraces a number of principles that align its provisions with the SPS Agreement. The first of these principles is state sovereignty, which recognizes that countries have the right to use phytosanitary measures, including measures taken in emergency situations, to protect their territories and their citizens from phytosanitary threats from other states. The effect of this right is, however, tempered by other principles, such as the principle of necessity, which requires states to adopt restrictive measures only where they are necessary for phytosanitary protection; and the principle of minimal impact (also contained in the SPS Agreement), which requires restrictive measures to have the least possible impact on the international movement of people and goods (IPPC, art. VII.2). Another important principle is cooperation, which requires states to cooperate to prevent the spread and introduction of quarantine pests and to promote measures for their official control (art. VIII).

The principle of non-discrimination requires that phytosanitary measures be applied without discrimination between countries with the same

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6 International Plant Protection Convention (New Revised Text Approved by the FAO Conference at its 29th Session – November 1997).
7 See Part VII.
phytosanitary status. In the case of regulated pests within a country, measures are to be applied without discrimination between domestic and imported consignments. The principle of transparency requires countries to publish and disseminate phytosanitary prohibitions, restrictions and requirements and, on request, to make available the rationales for them (art. VII.2). The principle of emergency action permits countries in the face of a new or unexpected phytosanitary situation to take immediate emergency measures on the basis of a preliminary pest risk analysis. Such measures are to be temporary and the validity of their application in the long term is subject to a detailed pest risk analysis as soon as possible (art. VII.6).

The SPS Agreement identifies the IPPC as the organization responsible for international phytosanitary standard setting. The IPPC secretariat established its standard-setting programme in 1992. The first International Standards for Phytosanitary Measures (ISPMs) were approved by the FAO Conference in 1995. From 1998 to 2005 they were approved by the Interim Commission on Phytosanitary Measures, now the Commission on Phytosanitary Measures. Twenty-seven ISPMs have been approved to date.8

Like the SPS Agreement, the New Revised Text of the IPPC makes provision for contracting parties to provide technical assistance to other contracting parties, especially developing countries, with the objective of facilitating implementation of the IPPC and its standards (art. XX).

IV. ANIMAL HEALTH

The Office international des épizooties (OIE), or World Organization for Animal Health, is designated under the SPS Agreement as the standard-setting body for animal health. The OIE has three main objectives: (1) to inform governments of the occurrence and course of animal diseases and of ways to control disease outbreaks; (2) to coordinate international scientific research on the surveillance and control of animal diseases; and (3) to facilitate the harmonization of regulations pertaining to trade in animals and animal products.

OIE member countries, usually through their official veterinary services, are obligated to collect information on animal diseases extant in their territories, which the OIE then analyses and distributes in order to facilitate prevention

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8 See J. Vapnek and D. Manzella, supra note 3, p. 4.
and control elsewhere. The OIE also provides expertise and technical support to member countries requesting assistance with animal disease control and eradication operations, including for diseases transmissible to humans (zoonoses). In addition, the OIE develops standards for international trade in animals and animal products, again with the intention of preventing the transmission of animal diseases.

OIE member countries must immediately report outbreaks of certain diseases and also periodically report on the presence and distribution of those diseases. WTO member states are allowed to take zoosanitary measures, including import controls, based on that information. They are also expected to submit their national regulations, particularly those that apply to imports, to the OIE. States may apply different standards only where the importing country demonstrates scientifically that national animal health conditions require standards over and above those established by the OIE.

The OIE develops and updates normative documents, such as the Terrestrial Animal Health Code, the Manual of Standards for Diagnostic Test and Vaccines, the Aquatic Animal Health Code and the Diagnostic Manual for Aquatic Animal Diseases, all of which contain a list of definitions, disease notification criteria, procedures for international reporting of diseases, principles for import risk analysis and import and export procedures. The standards, guidelines and recommendations contained within these health codes apply to trade in animals, animal genetic material and animal products. WTO member countries can use the information in these documents to devise measures to protect against animal diseases without setting up unjustified trade barriers.\(^9\)

V. FOOD SAFETY\(^{10}\)

The Codex Alimentarius (Latin for "food code") and its organization, the Codex Alimentarius Commission (Codex), serve as a global reference point for consumers, producers and national food regulatory agencies on internationally adopted food standards, codes of practice and residue limits of pesticides and veterinary drugs. Codex is recognized by the SPS Agreement as the source of


\(^{10}\) This section draws from J. Vapnek and M. Spreij, *Perspectives and Guidelines on Food Legislation, with a New Model Food Law*, FAO Legislative Study No. 87, 2005, pp. 29–37.
international food safety standards. The Codex Alimentarius contains more
than 200 standards for individual foods or groups of foods.

The preparation of draft food standards and related texts, whether intended
for worldwide use, for a given region or for a select group of countries, takes
place in Codex committees. Membership in these committees is open to all
Codex member states, and international organizations may attend (as
observers) committee sessions that are of interest to them. Generally,
committees are financially maintained and hosted by member states. The two
types of Codex committees are Commodity Committees and General
Subject Committees.

Codex Commodity Committees are often referred to as vertical committees
because they develop standards that apply to aspects of specific foods or
classes of food. Such standards generally concern quality factors such as the
composition or presentation of certain products. The subject matters of the
Codex Commodity Committees range from fresh fruits and vegetables to
processed meat and poultry products. Currently, eleven such committees are
active or in recess. Some of these committees have completed their work and
have ceased operation for an unspecified period of time until there is the
need to call them back into service, while still others have remained active
for the purpose of reviewing standards in order to bring them in line with
current practice.

In recent years, there has been a shift in focus away from quality concerns
towards food safety and the protection of human health. Thus, within Codex
attention has turned to "horizontal" subjects – food hygiene, labelling,
additives and contaminants – which, unlike vertical standards, cut across
different types and classes of foods. As a result, the Codex General Subject
Committees have grown in responsibility and prominence. These
committees develop concepts and principles applicable to foods in general or
applicable to specific foods or groups of foods, reviewing provisions in
Codex commodity standards and developing recommendations pertaining to
consumer health and safety. Currently, there are nine such committees,
including the Committee on Food Additives and Contaminants, the
Committee on Food Hygiene and the Committee on Food Labelling.

In addition to the established committees, from time to time Codex,
following its rules of procedure, establishes ad hoc task forces to deal with
specific new problems and issues. At present, one ad hoc task force is in the
process of developing standards, guidelines and recommendations for foods derived from biotechnology. The ad hoc task forces function in the same manner as the Codex General Subject and Commodity Committees except that they are dissolved after the specified work is completed or when the time limit allocated for the work has expired.

In addition to its many food standards, the Codex Alimentarius contains advisory instruments such as guidelines, principles, recommendations and codes of practice, with the goal of improving compliance with Codex standards. The codes of hygienic practice provide guidance on the production of food that is safe and suitable for consumption, while the codes of technological practice aim to ensure that the processing, transport and storage of food are carried out such that consumers receive end products that are wholesome and of the requisite quality. Many of these Codex instruments have been revised and updated over the years. For example, the Recommended International Code of Practice on General Principles of Food Hygiene, which is one of the most widely used Codex texts applying to all foods, has been revised four times since its adoption. During its recent revisions, the concept of risk analysis, as well as management tools such as the Hazard Analysis and Critical Control Point (HACCP) system, were included to emphasize the food chain approach, from primary production through to final consumption, highlighting the key hygiene controls required at each stage.

New instruments have been prepared over the last decade as well. For example, Guidelines for the Production, Processing, Labelling and Marketing of Organically Produced Foods were developed in 1999 in light of the growing production of and international trade in organically produced food, with a view to facilitating trade and preventing misleading claims. There are also several noteworthy initiatives in the area of biosafety. For example, the ad hoc Intergovernmental Task Force on Foods Derived from Biotechnology developed Principles for the Risk Analysis of Foods Derived from Modern Biotechnology, Guidelines for the Conduct of Food Safety Assessment of Foods Derived from Recombinant-DNA Plants and Guidelines for the Conduct of Food Safety Assessment of Foods Derived from Recombinant-DNA Microorganisms, which were adopted as official Guidelines at the 26th Session of Codex in July 2003.

More than forty years after its creation, the Codex Alimentarius has become the authoritative collection of internationally adopted food standards
covering all the principal foods traded internationally, whether processed, semi-processed or raw. The Codex Alimentarius is also supplemented by the many maximum residue limits established for pesticides in foods and animal feeds, residue levels for veterinary drugs in foods of animal origin and acceptable levels of food additives and contaminants.

VI. INVASIVE ALIEN SPECIES AND BIODIVERSITY

Invasive alien species (IAS) are species introduced deliberately or unintentionally outside their natural habitats into habitats where they have the ability to establish themselves, invade, out-compete natives and take over their new environment. IAS are relevant to Biosecurity because they have the ability to affect the human, animal and plant life and health of their new habitats. Moreover, they are of interest since the modern vision of Biosecurity includes a concern for the preservation of the environment and prevention of loss of biological diversity.

The most important international instrument in this sector is the Convention on Biological Diversity (CBD),\textsuperscript{11} which has three main objectives: (1) conserving biological diversity, (2) promoting the sustainable use of its components and (3) encouraging equitable sharing of the benefits arising out of the utilization of genetic resources (art. 1). Biological diversity is defined in the CBD as "the variability among living organisms from all sources, including, \textit{inter alia}, terrestrial, marine and other aquatic ecosystems and the ecological complexes of which they are part; this includes diversity within species, between species and ecosystems" (art. 2).

Article 8 of the CBD deals directly with IAS, providing that member states must prevent the introduction of, control or eradicate any IAS which threaten ecosystems, habitats or species. IAS have also been addressed by the CBD’s Conference of the Parties, which approved Guiding Principles for the Prevention, Introduction and Mitigation of Impacts of Alien Species that Threatens Ecosystems, Habitats or Species.\textsuperscript{12} These guidelines endorse a systematic approach to the control of IAS along the following lines:

- priority attention should be given to preventing the entry of potential IAS, both between and within states;

\textsuperscript{11} Convention on Biological Diversity (1992).
\textsuperscript{12} CBD, \textit{Guiding Principles for the Prevention, Introduction and Mitigation of Impacts of Alien Species that Threaten Ecosystems, Habitats or Species}, 2003.
• if entry has already taken place, actions should be undertaken to prevent the establishment and spread of alien species;
• the preferred response is eradication at the earliest possible stage; and
• if eradication is not feasible or cost-effective, containment and long-term control measures should be considered.

Other CBD provisions are pertinent to the conservation of the environment and biological diversity and are therefore also relevant to Biosecurity. Member states are required to ensure that activities within their jurisdiction or control do not cause damage to the environment of other states or areas beyond the limits of national jurisdiction (art. 3). If there is imminent danger or damage to biodiversity either originating under a member state’s jurisdiction or within its control outside its jurisdiction, that state must immediately notify potentially affected states of such danger or damage, as well as initiate action to prevent or minimize such danger or damage (art. 14).

Biotechnology is defined under the CBD as any technological application that uses, inter alia, living organisms to make or modify products or processes for a specific use (art. 2). Member countries are required to "take all practicable measures" that would give priority access to the results or benefits that come from biotechnologies based on genetic resources (art. 19(2)). Parties were mandated to consider the need for, and the modalities of, a protocol setting out appropriate procedures for the safe transfer, handling and use of any living modified organism that may have adverse environmental impacts that could affect the conservation and sustainable use of biological diversity (art. 19(3)). This obligation was fulfilled by the adoption of the Cartagena Protocol on Biosafety (Cartagena Protocol).

VII. LIVING MODIFIED ORGANISMS

The Cartagena Protocol was adopted by the Conference of the Parties (COP) of the CBD on 29 January 2000 and came into force on 11 September 2003. The objective of the protocol is "to contribute to ensuring an adequate level of protection in the field of the safe transfer, handling and use of living modified
organisms resulting from modern biotechnology that may have adverse effects on the conservation and sustainable use of biological diversity, taking also into account risks to human health, and specifically focusing on transboundary movements (art. 1).

Although the protocol is basically an environmental instrument, it does include within its objectives protection against the possible impacts of living modified organisms (LMOs) on human health. The protocol recognizes that there are intrinsic risks associated with LMOs – both to the environment and human health – and promotes biosecurity by setting the rules for the safe transfer, handling and use of LMOs, focusing on the transboundary movement of LMOs intended for the release into the environment. Among other things, the protocol requires that shipments of LMOs intended for intentional introduction into the environment be accompanied by documentation clearly stating that the shipment contains LMOs (art. 18).

The Cartagena Protocol provides for an "advance informed agreement" (AIA) between the exporting state and the importing state. The AIA involves a notification in writing by the exporting state before it exports a consignment of LMOs to the importing state. Crucially, upon receipt of this notification, the importing state must indicate whether its own regulations or those provided in the protocol will have to be followed with regard to the importation (art. 9.2(c)). The importing state shall make a decision on the importation within a prescribed time frame (art. 10.3) and has a right to refuse entry of the consignment of LMOs based on risk assessment (art. 10.1); the parameters of these risk assessment procedures are contained in Annex III to the protocol. Failure to acknowledge receipt of notification does not imply that the movement of the LMOs is permitted (art. 9.4). Similarly, failure to communicate the decision within the prescribed time frame does not imply any consent to the movement of LMOs (art. 10.5).

The protocol provides that in case of scientific uncertainty regarding the potential adverse effects of the LMOs in question, the importing state can

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13 "Living modified organism" is defined as "any living organism that possesses a novel combination of genetic material obtained through the use of modern biotechnology" (art. 3(g)). The term is wider than genetically modified organism (GMO) as it does not require the insertion of genetic material. However, in many countries the term GMO is used to cover LMOs. See R. Mackenzie, F. Burhenn-Guilmin, A. La Viña and J. Werksman, An Explanatory Guide to the Cartagena Protocol on Biosafety, IUCN, 2003, p. 46.
still decide against the import in order to prevent the possible adverse effects (art. 10.6). Thus the protocol embraces the precautionary principle.

It is important to note that the AIA procedure only applies to those LMOs intended for intentional introduction into the environment, and not to:

1. LMOs identified in a decision of the COP as not likely to have adverse effects on biodiversity conservation and sustainable use,
2. LMOs in transit,
3. LMOs for contained use,
4. Pharmaceuticals that are intended for human use,
5. LMOs intended for direct use as food or feed or for processing (art. 5).

With regard to the last category, governments that approve LMOs for domestic use or for import shall communicate this decision and related information to the Biosafety Clearing-House (BCH) Mechanism.

The BCH is an information exchange platform established in the protocol and designed to facilitate the exchange of scientific, technical, environmental and legal information and experience on LMOs. Member states of the protocol are required to make available to the BCH any information regarding their national biosafety situation, including existing laws, regulations and guidelines for implementation of the protocol, information required for the AIA, any bilateral, regional and multilateral agreements and arrangements as well as summaries of risk assessments and final decisions (art. 20).

VIII. OTHER INSTRUMENTS

Biosecurity covers a wide range of sectors, including the food safety aspects of food production and fisheries. Hence, sectoral international instruments in those areas, binding and non-binding, are relevant to Biosecurity for the purposes of food and agriculture and can be said to form part of the international Biosecurity framework. An illustrative (but not exhaustive) list would include the following instruments:

- Rotterdam Convention on the Prior Informed Consent Procedure for Certain Hazardous Chemicals and Pesticides in International Trade;
- Convention on Persistent Organic Pollutants;
- FAO International Code of Conduct on the Use and Distribution of Pesticides;
- Biological and Toxin Weapons Convention;

14 See Biosecurity in Food and Agriculture, FAO Committee on Agriculture, 17th Session, 31 March–4 April 2003, Rome.
International Framework

- FAO International Code of Conduct on Responsible Fisheries;
- Ramsar Convention on Wetlands;
- Protocol to the Antarctic Treaty on Environmental Protection;
- Bonn Convention on the Conservation of Migratory Species of Wild Animals;
- Global Programme of Action for the Protection of the Marine Environment from Land-based Activities;
- United Nations Framework Convention on Climate Change;

In the discussion that follows, these international instruments are grouped according to those aspects of Biosecurity to which they are most relevant, using the same general Biosecurity categories as above: (1) food safety and plant and animal life and health and (2) IAS and LMOs.

Food Safety, and Plant and Animal Life and Health

(a) Rotterdam Convention

The Rotterdam Convention on the Prior Informed Consent Procedure for Certain Hazardous Chemicals and Pesticides in International Trade\(^{15}\) seeks to protect human health and the environment from the possible risks resulting from trade in highly dangerous pesticides and chemicals by creating legally binding obligations for the implementation of a prior informed consent procedure by importing countries. As pesticides and chemicals pose risks to both the environment and food safety, the convention is relevant to Biosecurity.

(b) POPs Convention

The Stockholm Convention on Persistent Organic Pollutants\(^{16}\) (POPs) is intended to eliminate or restrict the production, use or release of a dozen POPs including pesticides, industrial chemicals and hazardous by-products of combustion. Like the Rotterdam Convention, the POPs Convention aims to protect human health and the environment from substances that are toxic to humans and wildlife, and thus has Biosecurity implications.

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(c) FAO International Code of Conduct on Pesticides

The FAO International Code of Conduct on the Distribution and Use of Pesticides\textsuperscript{17} sets forth voluntary standards for governments and the private sector on pesticides management. The code embodies principles such as risk reduction and support for sustainable agricultural development. As the code aims to protect human and environmental health, it is relevant to \textit{Biosecurity}.

(d) Biological and Toxin Weapons Convention

The so-called Biological and Toxin Weapons Convention\textsuperscript{18} prohibits the development, production and stockpiling of biological and toxin weapons. The convention is relevant to \textit{Biosecurity} from the unique perspective of biological warfare, which may specifically target plants or crops.

(e) Code of Conduct for Responsible Fisheries

The FAO Code of Conduct for Responsible Fisheries\textsuperscript{19} is widely recognized by governments and non-governmental organizations as the global standard for sustainable fisheries and aquaculture. It sets out principles and international standards of behaviour for responsible practices with a view to ensuring the effective conservation, management and development of living aquatic resources, with due respect for ecosystems and biodiversity. It is the basis for reviewing and revising national fisheries legislation, which may include provisions on the prevention of fish diseases.\textsuperscript{20}

As \textit{Biosecurity} includes fisheries, the aquatic animal health and food safety provisions of the code are important elements of the international framework for \textit{Biosecurity}. For instance, the code mandates states to promote effective farm and fish health management practices favouring hygienic measures and vaccines (art. 9.4.4). States should also regulate the use of chemical inputs in aquaculture which are hazardous to human health and the environment (art. 9.4.5).

\textsuperscript{17} International Code of Conduct on the Distribution and Use of Pesticides (2002).
\textsuperscript{20} See FAO, \textit{Law and Sustainable Development Since Rio, Legal Trends in Agriculture and Natural Resources Management}, FAO Legislative Study No. 73, 2002, p. 194.
Invasive Alien Species and Living Modified Organisms

(a) Ramsar Convention

The Ramsar Convention on Wetlands (Ramsar Convention) is an international treaty that provides the framework for national action and international cooperation for the conservation and wise use of wetlands and their resources. Although the convention’s text itself does not mention invasive alien species (IAS), contracting parties addressed the topic of "Invasive Species and Wetlands" in Resolution VII/14. The resolution calls upon member states to address the impact of IAS on wetlands within their jurisdictions and to identify methods of control and solutions for combating IAS.

(b) Protocol on Environmental Protection to the Antarctic Treaty

Annex II to the Protocol on Environmental Protection to the Antarctic Treaty has provisions connected to the international biosecurity framework. For example, article 4 imposes upon member states the requirement that no non-native species can be introduced into Antarctica, except with a permit (art. 4.1).

(c) Bonn Convention

The Convention on the Conservation of Migratory Species of Wild Animals (Bonn Convention) requires its member states to endeavour "to the extent feasible and appropriate to prevent, reduce or control factors that are endangering or are likely to further endanger certain species, including strictly controlling the introduction of, or controlling or eliminating, already introduced exotic species" (art. III(4)(c)). Any action taken to implement those provisions may be part of an overall IAS risk management programme.

22 Resolutions of the San Jose Conference, Resolution VII.14 on Invasive Alien Species and Wetlands.
(d) Global Programme of Action for the Protection of the Marine Environment from Land-based Activities

The Global Programme of Action for the Protection of the Marine Environment from Land-based Activities\(^25\) was convened by the Executive Director of the United Nations Environment Programme (UNEP). One of its goals is to prevent the introduction of alien species known to have serious effects upon marine ecosystem integrity. Its overall aim is to help facilitate the preservation of the marine environment through international legal obligations such as the United Nations Convention on the Law of the Sea (UNCLOS) and Agenda 21.

(e) UNFCCC

The United Nations Framework Convention on Climate Change\(^26\) (UNFCCC) has the aim of stabilizing (and eventually reducing) greenhouse gas concentrations in the atmosphere so as to prevent dangerous anthropogenic interference with the climate system. The convention and its Kyoto Protocol are of relevance to Biosecurity in that they attempt to prevent climate effects which will have an effect on biodiversity and on the movement of IAS.

(f) UNCLOS

Pursuant to UNCLOS,\(^27\) member states must take all measures necessary to prevent, reduce and control "the intentional or accidental introduction of species, alien or new, to a particular part of the marine environment, which may cause significant and harmful changes thereto" (art. 196). These provisions can be interpreted to support, for example, risk assessment for genetically modified organisms prior to their release into the marine environment.

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IX. CONCLUSION

Biosecurity covers a number of subjects, each with its own intricacies. The international regulatory instruments introduced here also deal with the components of Biosecurity from sectoral perspectives. The task for national governments will be to identify conventions and international agreements which it is bound to or desires to follow, and assess its national legislation for conformity with those agreements. The main international instruments relevant to Biosecurity described in Parts III to VII of this chapter should be the starting point for the analysis.

Identifying other international agreements relevant to the adoption of a Biosecurity approach, such as those set out in Part VIII, may be a more challenging task. The key for national lawmakers will be to examine the constellation of potentially relevant international instruments from the perspective of whether they have an impact on the management of risks to food and agriculture. In this era of burgeoning international trade, national governments must carry out a delicate balance: implementing their international obligations and aligning their national laws with these international obligations, while structuring their national legal and institutional frameworks in a manner most conducive to the protection of their natural resources for food and agriculture.
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# REVIEW OF PILOT CASE STUDIES

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I. LEGISLATION OVERVIEW

National legal consultants in Ethiopia, Ghana, India, Kenya, Uganda, and Viet Nam examined a range of national legislative instruments in the various sectors of *Biosecurity*. The instruments included parliamentary-level and subsidiary legislation (such as regulations and ministerial orders), and consisted of legislation in force as well as draft instruments under consideration.

This section sets out a list of the legislation reviewed in each country\(^1\) while the subsequent sections contain a gap analysis of each of the five sectors of the legislative framework for *Biosecurity*: food safety, plant health, animal health, invasive alien species and biosafety. The chapter concludes with a review of the institutional set-up for *Biosecurity* controls in each of the six countries.

The Ethiopia study was based on the following pieces of legislation:

<table>
<thead>
<tr>
<th>SECTOR</th>
<th>LEGISLATION</th>
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<tbody>
<tr>
<td>Food Safety</td>
<td>Quality and Standards Authority of Ethiopia Proclamation No. 413/2004;</td>
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<tr>
<td></td>
<td>Re-establishment and Modernization of Customs Authority (Amendment) Proclamation, No. 368/2003;</td>
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<td></td>
<td>Public Health Proclamation, No. 200/2002;</td>
</tr>
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<td></td>
<td>Drug Administration and Control Authority Proclamation No. 176/1999;</td>
</tr>
<tr>
<td></td>
<td>Ethiopian Health and Nutrition Research Institute Establishment, Council of Ministers Regulations No. 4/1996;</td>
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<tr>
<td></td>
<td>Ministry of Health Proclamation No. 4/1995.</td>
</tr>
<tr>
<td>Animal Health</td>
<td>Animal Diseases Prevention and Control Proclamation No. 267/2002;</td>
</tr>
<tr>
<td></td>
<td>Draft Regulation for Animal Diseases Prevention and Control, 2000;</td>
</tr>
<tr>
<td></td>
<td>Draft Regulation for Controlling Movement of Animals and Transportation of Animal Products and By-products, 2000;</td>
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<tr>
<td></td>
<td>Draft Regulations to Provide for the Registration and Licensing of Animal Health Professionals, 2000;</td>
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</tbody>
</table>

\(^1\) A detailed description of the content of the various legislative instruments in the six countries can be found in Chapters 5–10.
**SECTOR** | **LEGISLATION**
---|---


**Biosafety** | Draft National Biosafety Framework, 2000.2

In Ghana, the following instruments were analysed:

**SECTOR** | **LEGISLATION**
---|---

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2 The National Biosafety Framework is a policy document containing an outline of the biosafety legislation to be drafted.
<table>
<thead>
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<th>SECTOR</th>
<th>LEGISLATION</th>
</tr>
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<tr>
<td>Invasive Alien Species</td>
<td>Wetland Management (Ramsar Sites) Regulations, 1999 (LI 1659).</td>
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</table>

In India, the country study is based on the following legislation:

<table>
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<tr>
<th>SECTOR</th>
<th>LEGISLATION</th>
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<tbody>
<tr>
<td>Animal Health</td>
<td>Livestock Importation Act, 1989; Wild Life Protection Act, 1972</td>
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</table>
In Kenya, the following legislation was analysed:

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<tr>
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<th>LEGISLATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Animal Health</td>
<td>Animal Diseases Act, Chapter 364, 1972, as revised in 1989; Cattle Cleansing Act, Chapter 359, 1937; Crop Production and Livestock Act, Chapter 321, 1926, as last amended in 1968.</td>
</tr>
<tr>
<td>Invasive Alien Species</td>
<td>Environmental Management and Coordination Act, No. 8 of 1999.</td>
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</tbody>
</table>
The national legal consultant reviewed the following legislation in Uganda:

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<th>SECTOR</th>
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<tr>
<td>Food Safety</td>
<td>Uganda National Bureau of Standards Act, Chapter 327, No. 1, 1983;</td>
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<td>Public Health Act, Chapter 281, 1964;</td>
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<td>Food and Drugs Act, Chapter 278, 1964.</td>
</tr>
<tr>
<td>Animal Health</td>
<td>Animal Breeding Act, 2001;</td>
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<td>Uganda Wildlife Act, Chapter 200, 2000;</td>
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<td></td>
<td>Veterinary Surgeons Act, Chapter 277, 1966;</td>
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<td>Cattle Traders Act, Chapter 43, 1964;</td>
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<td>Animal Diseases Act, Chapter 38, 1964.</td>
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<td>Plant Health</td>
<td>Draft Plant Protection Bill, 2006;</td>
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<td>Agricultural Seeds and Plant Act, Chapter 28, 1994;</td>
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<td>Plant Protection Act, Chapter 31, 1976.</td>
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<td>Biosafety</td>
<td>Draft National Biotechnology and Biosafety Policy, 2006;</td>
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The Viet Nam study reviewed the following legislation:

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<th>SECTOR</th>
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<td>Food Safety</td>
<td>Decision No. 21/2007/QD-BYT of the Ministry of Health on Health Measures in Food Manufacturing Sites;</td>
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<td>Decree No. 21/2006/ND-CP of the Vietnamese Government on the trade in and use of nutrition products for children;</td>
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<td>Decision No. 43/2006/QD-TTg of the Prime Minister on the National Action Plan on Hygiene and Food Safety to 2010;</td>
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<td>Decision No. 117/2000/QD- BNN-BVTV on the Quarantine Pest List;</td>
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<td>Decision No. 70/1998/ QD-BNN-KHCN on Procedures for Fumigation;</td>
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II. FOOD SAFETY

2.1. Ethiopia

The Government of Ethiopia issued Public Health Proclamation No. 200 in 2002. In the proclamation, the Ministry of Health is given general powers on public health matters, which include food safety. The proclamation broadly defines food as "any substance whether processed, semi-processed or raw which is intended for human consumption and includes drinks, chewing gum, and/or treatment of food, not including tobacco, cosmetics or substances used only as drugs".3

The proclamation establishes an advisory board at the federal level and regional health bureaux for the purpose of advising the appropriate health

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3 The definition mainly tracks the Codex definition of food.
authority (the Ministry of Health) on the implementation of the proclamation. The proclamation sets forth general prohibitions of manufacture, import or sale of food not in compliance with national quality standards. Draft Food Safety Regulations detailing food safety controls have been elaborated and are awaiting endorsement by the Council of Ministers. Most regional health bureaux, under the Public Health Proclamation, have enacted regulations that fit their regional contexts.

The Meat Inspection Amendment Proclamation (No. 81/1976) and the Animal Diseases Prevention Control Proclamation (No. 267/2002) provide for the control of slaughterhouses and establishments as well as the safety of meat and meat products. These instruments are implemented by the Animal and Fisheries Resources Development and Regulatory Department of the Ministry of Agriculture.

By virtue of the national standard-setting mandate contained in Proclamation No. 413/2004, the Quality Standards Authority of Ethiopia (QSAE), which operates under the Ministry of Trade and Industry, has developed about 450 food-related standards, most of which have been translated into technical regulations. Currently, about 60 percent of the QSAE-approved standards fall under the category "Agriculture and Food Technology".

A number of institutions are assigned, via the proclamations establishing them, to undertake food safety inspections in the country. These include the Ministry of Health, the QSAE, the Ethiopia Health and Nutrition Research Institute and the Customs Authority. To strengthen collaboration, the existing Ethiopian Technical Committee has established the National Food Safety Council, a consultative body whose members are drawn from regulatory bodies, research institutes, industry, consumers and institutions of higher learning involved in food safety.

2.2. Ghana

In Ghana, the 1992 Food and Drugs Law (FDL) regulates the manufacture, importation, exportation, distribution, use and advertisement of foods, drugs, cosmetics, chemical substances and medical devices. It contains prohibitions against the sale of unwholesome, poisonous and adulterated foods and it prescribes standards for foods.
The FDL establishes an administrative authority, the Food and Drugs Board (FDB), under the control and supervision of the ministry responsible for health. The composition of the board draws from relevant departments and agencies of state and the private sector. The functions of the FDB as set out in Section 28 of the FDL include advising the ministry on all matters relating to the administration and implementation of the FDL.

A major defect of the FDL from the standpoint of Biosecurity is that it is void of any reference to international standards that should guide the FDB in the discharge of its duties. Schedule I of the FDL, which is linked to other national legislation on standard setting, makes reference to the publications of certain international bodies but omits the Codex Alimentarius Commission. Reportedly, a draft Food Bill was developed in 2006 to address this gap.

Authorized officers of the FDB have wide enforcement powers under the FDL for purposes of entering premises, opening and examining food receptacles and books and seizing and destroying unwholesome, poisonous or adulterated foods. Nonetheless, there is a gap in the legislation regarding meat inspection.

Under the FDL, both the FDB and the district/metropolitan assemblies have statutory functions in meat inspection. The meat inspection function has been exercised by public health officers by virtue of previous and current legislation on local government. Unlike these officials, whose mandate is expressly provided for in legislation, no specific mandate is accorded to veterinary officers of the Directorate of Veterinary Services (DVS) of the Ministry of Food and Agriculture in the area of meat inspection.

A revised draft Meat Inspection Bill was prepared in 2004 to divest public health officers of these functions and vest them in the DVS. The draft also makes provision for the appointment and qualifications of "veterinary inspectors". These include qualified and registered veterinarians and any other veterinary personnel appointed as inspectors pursuant to the law.

The Animals (Control of Importation) Ordinance, Diseases of Animals Act, 1961, a colonial statute still in force, bans the importation of animals into the country unless they are certified by a veterinary authority as free from disease. The ordinance is outdated and could be repealed by the draft Meat
Inspection Bill, which deals with importation and exportation of meat, meat products and animals.

Even though the appropriate international bodies have prescribed standards for inspections and the importation and exportation of meat products, there is no express reference to the OIE in the current legislation. Either the parent enactment could make a reference to these standards, or regulations could be passed incorporating them.

The Ghana Standards Board (GSB) is a statutory body that was established by the Standards Decree, 1967, and re-established by a new decree in 1973. The 1967 Decree grants the GSB a wide range of functions and powers on standard setting, implementation and enforcement. Standards cover the sale or manufacture of goods in the national interest as well as in the interest of public health and safety.

Another piece of legislation, the 1979 GSB Decree, added two specific functions to be exercised by the GSB in relation to food, namely: (1) prohibiting the sale or manufacture of foods, in the national interest; and, (2) prohibiting the importation into Ghana of foods which have not been certified by the GSB as compliant with its standards. Both the GSB and the FDB have statutory functions in the area of sale, manufacture, exportation and importation of food, and this has become a source of overlap and conflict between the two boards.

The 2006 draft Standards Bill establishes a National Standards Authority (NSA) as a body corporate. The bill re-enacts the provisions of the GSB decree and transfers the functions of the GSB to the NSA. The specifications for standards prescribed by law include "international or other overseas specifications", without explicit reference to WTO standards.

The draft bill, however, does not satisfactorily address the thorny issue of the NSA functions in the area of food vis-à-vis the FDB. Section 3(2)(c) of the draft bill states that if it is within the national interest the NSA is authorized to prohibit the sale or manufacture of any kind of goods. The NSA also has the power to prohibit the importation of goods that have not been certified as complying with the standards, and the definition of "goods" is wide enough to encompass food. Hence the draft bill in its present form conflicts with the mandate of the FDB which, as noted, has been established to control the manufacture, importation, exportation, distribution and use of food.
2.3. India

The Indian national case study notes that the SPS Agreement has prompted substantial changes, and not only in the food laws. At present, food safety legislation is still disparate, with several subordinate rules, regulations and orders having been enacted to deal with contingencies as they arose. The operative legislation, namely, the 1954 Prevention of Food Adulteration (PFA) Act, seeks to test only end products and does not foster the adoption of Hazard Analysis and Critical Control Point (HACCP) principles throughout the food chain.

The state governments and the union territories are responsible for monitoring and implementing the provisions of the PFA Act and the PFA Rules, 1955. The latter were adopted by the Ministry of Health and prescribe maximum tolerance limits for pesticides and heavy metals in food products. The Directorate General of Health Services in the Ministry of Health and Family Welfare, which is the Codex Contact Point, is currently working to integrate Codex standards into the legislation.

The Ministry of Food Processing Industries (MOFPI) is in charge of the implementation of a number of food safety and quality provisions. For example, the Fruit Products Order of 1955 promulgated under Section 3 of the Essential Commodities Act of 1955 prescribes minimum norms for sanitary and hygienic conditions of manufacturing premises and also lays down product standards. MOFPI is closely associated with the Codex Contact Point in the Ministry of Health.

The Food Safety and Standards Act (FSSA), 2006, which seeks to consolidate the many pieces of legislation into one combined whole, is a serious attempt at implementing a food chain approach, promoting a continuous series of controls from the farm to the table. However, the FSSA excludes from its purview plants prior to harvesting and animal feed and hence does not control the contamination of food from pesticides and antibiotics at source. The FSSA does however establish a Food Safety and Standards Authority.

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4 India is a federal republic which comprises 28 states and seven union territories.
5 See Part VII on the institutional set-up.
In India, international standards, guidelines and recommendations are increasingly used to guide domestic as well as international trade in food. The Codex HACCP and food hygiene standards have been adopted by the Bureau of Indian Standards (BIS), an autonomous statutory body set up by the Bureau of Indian Standards Act, 1986. The BIS comprises members representing industry, consumer organizations, scientific and research institutes, technical institutions, central ministries, state governments and members of parliament. It provides for quality certifications, including food hygiene.

Inspection and certification for export are regulated under the Export (Quality Control and Inspection) Act of 1963. The Export Inspection Council (EIC) is the official certification body for exports operating under the act. Notably, the EIC is developing equivalence agreements on conformity assessment with its major trade partners. It is also developing standards for exports based mainly on Codex standards, but it also recognizes that an importing country may impose stiffer requirements.

Imported food is inspected at the ports of entry by personnel of the Collectorate of Customs. The Government of India through its various departments – Commerce, Health, Revenue and the Directorate General of Foreign Trade – is taking steps to streamline the inspection of imported food.

2.4. Kenya

In Kenya, the Food, Drugs and Chemical Substances Act (Chapter 254, 1970) makes provision for the prevention of adulteration of food, drugs and chemical substances. Foods for which there are prescribed standards must conform to such standards. Subsidiary legislation under the act makes provisions for food hygiene, and has addressed the issues of food labelling, additives and standards.

Meat control is also the subject of specific legislation. The Meat Control Act (Chapter 356, 1973) provides standards for slaughterhouses; storage and transportation of meat and meat products intended for human consumption; meat processing establishments; and import and export control over meat and meat products. Regulations specify standards to be observed in meat production as well as methods of packaging, labelling and transport. The Ministry of Agriculture implements both the Food, Drugs and Chemical Substances Act and the Meat Control Act.
The Standards Act (Chapter 496, 1974, as amended) is the main legislation for standards formulation and implementation in Kenya. Section 3 establishes the Kenya Bureau of Standards, whose functions inter alia are to make arrangements and provide facilities for the examination and testing of "commodities and any material substance from or with which they may be manufactured, produced, processed or treated". Those provisions are broad enough to cover food. The minister in charge of trade is empowered under the act to appoint inspectors who are mandated to, among other things, inspect and take samples of any commodity or any related material or substance.

Food is also regulated under public health legislation. The Public Health Act (Chapter 242, 1921, as amended) establishes a Central Board of Health, which is empowered to advise the Minister of Health on all matters affecting health. The act contains provisions that ensure the protection of foodstuffs intended for human consumption. Another significant provision on food safety is the requirement that local authorities ensure that water supplies, food and milk are wholesome.

2.5. Uganda

The Food and Drugs Act (Chapter 278, 1964) is the main piece of legislation on food safety in Uganda. The act makes provision for the prevention of adulteration of food, which is defined to include drink, chewing gum and other products of like use or nature, and articles and substances used as ingredients in the preparation of food or drink or of such products. It excludes water, live animals or birds, animal fodder or feed and substances used only as drugs (sect. 1). The act proscribes the use of any ingredient in the preparation of food sold for human consumption that would render the food injurious to human health (sect. 2) and prohibits false labelling or advertisement of food (sect. 5). Food in transit in Uganda may be examined by an authorized officer (sect. 9).

An authorized officer means a person authorized by the Minister of Health or a local authority with the approval of the minister. For the purposes of taking samples, an authorized person includes a police officer of or above the rank of inspector authorized to take samples. A veterinary surgeon registered under the Veterinary Surgeons Act (Chapter 277, 1966), in the service of the government or of a local authority, is deemed to be an authorized officer for the purposes of the inspection of animals intended for
slaughter and the examination and seizure of meat unfit for human consumption. A medical officer, a health inspector or a person having such qualifications as may be prescribed may undertake certain functions of the veterinary surgeon.

The Food Hygiene Advisory Committee is established under the act to advise the minister on any questions relating to the act that the minister may refer to it for its consideration.

The Uganda National Bureau of Standards (UNBS) Act (Chapter 327, 1983) is of relevance to food safety in that it establishes the UNBS under the general supervision of the minister responsible for commerce. The functions of the UNBS include the formulation of national standards and specifications for commodities, including food, as well as standards enforcement to protect consumers against harmful ingredients and dangerous components of commodities.

2.6. Viet Nam

In Viet Nam, food safety is mainly regulated by the 2003 Ordinance on Hygiene and Food Safety. The ordinance includes provisions to ensure hygiene and food safety in food production and trade as well as prevention and control of food poisoning and food-borne diseases. The ordinance establishes that individuals and legal entities manufacturing and trading in food must comply with three sets of regulations: (1) safety regulations on infrastructure, including facilities, water supply systems and wastewater treatment; (2) regulations on equipment, such as for processing, storage and transportation; and (3) regulations on personnel, such as employees’ health and knowledge of hygiene and food safety principles. National standards of food hygiene and safety are established by the Ministry of Science and Technology in cooperation with the Ministry of Health.

Provisions on food export and import require import and export enterprises to obtain an authorization from the Ministry of Health, which certifies that the enterprises have an adequate food safety management infrastructure. In cases where the requirements are not met, food may be seized and disposed of.

In the area of prevention and control of food poisoning and food-borne diseases, the People’s Committees at different territorial levels, the Ministry of Agriculture and Rural Development, the Ministry of Industry and the
Ministry of Health are all assigned a number of responsibilities. These range from the implementation of good manufacturing practices to the implementation and enforcement of food safety and hygiene standards and food safety emergency management.

Other laws, such as the 2000 Law on People's Health and the 1999 Ordinance on Consumer Protection, state the general duty of individuals and legal entities to follow food safety and hygiene standards.

III. PLANT HEALTH

3.1. Ethiopia

The Council of Ministers Regulations No. 4/1995 give the Ministry of Agriculture and Rural Development the general mandate for plant health. The ministry is made responsible for plant quarantine to prevent the spread of plant pests and to regulate the movement of plants, plant products or other articles into or from a specified area.

The ministry is further empowered to restrict the importation of certain plants and plant products by requiring import permits and phytosanitary certificates duly issued by the plant protection authorities of the exporting countries. The ministry has the responsibility for issuing phytosanitary certificates for export of plants and plant products.

The Plant Quarantine Regulations (Council of Ministers Regulations No. 4/1992) elaborate detailed provisions on import and export. The regulations prescribe that any plants or other articles, premises or conveyances found to be infected shall be treated or destroyed, as the case may be. Quarantine controls and documentary verification of phytosanitary certificates on all imported plants are required. Some plant species are prohibited from entering the country. The regulations also provide for the declaration of quarantine areas and the adoption of subsequent control measures.

3.2. Ghana

In Ghana, the major piece of legislation governing plants and plant protection is the Prevention and Control of Pests and Diseases of Plants Act, 1965. The act regulates the prevention of plant pests and also governs plant quarantine. It confers the general mandate for plant protection on the
Ministry of Food and Agriculture and provides for the appointment of plant quarantine officers. At the borders, officials of the Plant Protection and Regulatory Services Department carry out inspections on all imported plants and plant materials in accordance with the act.

The legal regime for plant protection is outdated, and the legislation does not measure up to IPPC standards. The shortcomings include the absence of provisions on the designation of a national plant protection organization, on risk analysis and on the exportation of plants, as well as insufficient financial penalties for violations.

Draft legislation was prepared with the assistance of FAO in the mid-1990s but has not yet been enacted. It is also somewhat out of date given the coming into force of the New Revised Text of the IPPC in 1997.

3.3. India

The Destructive Insects and Pests Act, 1914, continues to regulate the introduction and movement of any insect, fungus or pest which would be destructive to crops and crops only (not to areas such as forests). It has gone through several amendments over the years.

The act does not regulate the export and certification of plants and plant products. The enactment of the Plant Quarantine Bill, 2004, would repeal it and provide a comprehensive regulatory framework for quarantine pests. The bill establishes the Plant Quarantine Authority of India as the national plant protection organization, thus meeting India’s obligation under the IPPC.

With regard to imports, the Plant Quarantine (Regulation of Import into India) Order, 2003, supplements the 1914 Act. The order classifies plants and plant products for import as: (a) prohibited; (b) restricted (i.e. subject to a special authorization regime in addition to ordinary import conditions); (c) requiring additional declarations and other import conditions; and (d) requiring phytosanitary certification for processing and industrial production. The central government, through the Joint-Secretary in charge of plant protection in the Department of Agriculture and Cooperation, can relax any of the conditions of the order for public interest reasons. The power to relax conditions on import permits and phytosanitary certificates has been delegated to officers in charge of plant quarantine stations.
There is generally a lack of enforcement of the existing legislation and an inability to follow the letter of the law (for instance, on phytosanitary certification). However, the current framework is in broad compliance with international standards, and the frequent updating of the Plant Quarantine Order suggests that the concerned ministerial department is trying to protect domestic plant health adequately while at the same time pay due attention to international trade requirements.

3.4. Kenya

In Kenya, the main legislation on plant health is the Plant Protection Act (Chapter 324, 1962, as amended). The act makes provision for the prevention of the introduction and spread of diseases destructive to plants. The main regulatory agents under the act are the Minister of Agriculture and the inspectors, including the Director of Agriculture and any other persons authorized by the director to enforce the act.

In the Plant Protection Act, the Minister of Agriculture is given regulatory powers in relation to:

(a) phytosanitary inspection and certification for imports and exports;
(b) disinfection or treatment of any plant or article likely to infect any plant with a pest or disease;
(c) imports through specified ports or places of entry;
(d) post-entry quarantine; and
(e) the movement of plants or classes of plants likely to be infected with any pest or disease into or within any specified place or area.

Under the act, inspectors are mainly appointed from staff of the Kenya Plant Health Inspectorate Service (KEPHIS). KEPHIS is a parastatal agency under the Ministry of Agriculture that was established by ministerial order under the provisions of State Corporations Act (Chapter 446, 1986). KEPHIS is the SPS Enquiry Point. Its mandate is to:

(a) coordinate all matters relating to crop pests and disease control;
(b) establish service laboratories to monitor the quality and levels of toxic residues in plants as well as soils and produce;
(c) advise the Director of Agriculture on appropriate seeds and planting materials for import and export;
(d) undertake inspection, testing, certification, quarantine control, variety testing and description of seeds and planting materials;
(e) undertake grading and inspection of plants and plant produce at the ports of entry and exit;
(f) develop and implement standards on both imported and locally produced seeds;
(g) approve all import and export licences for plants and seeds issued by the ministry responsible for commerce and industry, before such importation or exportation is carried out; and
(h) establish posts at convenient locations for quarantine, inspection and quality control of fertilizers and seeds.

The Plant Protection Act is complemented by the Suppression of Noxious Weeds Act (Chapter 325, 1986), which provides that the Minister of Agriculture may, by notice in the gazette, declare a plant to be a noxious weed in any area. The inspectors, appointed by the Director of Agriculture, are granted powers of entry onto land for the purpose of ascertaining whether any noxious weed exists and, if so, to serve notice on the person in charge of the land. None of the legislation examined above refers to the IPPC or specifically to the mandate of the national plant protection organization.

3.5. Uganda

Uganda’s legislation on plant health reveals an outdated framework that ought to be aligned with international requirements if it is to facilitate agricultural imports and exports. The Plant Protection Act (Chapter 31, 1976) was originally passed as an ordinance in 1937. The act has limited provisions on the prevention of the introduction and spread of diseases destructive to plants. Of course, the definition section does not reflect the modern concepts of plant protection even as first defined in the IPPC of 1951.

The existing plant protection administration is undersized and does not allow for the delivery of an efficient service. The penalties set in the legislation are outdated and have no deterrent effect. The review of the act started in 2001 led to the drafting of the Plant Protection and Health Bill of 2003.

The 2003 Bill attempted to fill the gaps by establishing a Technical Committee to assist the Commissioner and the Minister of Agriculture in carrying out the functions outlined for the Department of Crop Protection of the Ministry of Agriculture. The penalties were reviewed and currency points introduced to
make the penalties more realistic. The definition section was expanded to include modern terminology, drawing on the Glossary of Phytosanitary Terms (ISPM No. 5). However, the cost recovery in the draft bill, proposed in particular to enable rapid response to epidemics of quarantine importance, was not included in the final text. The 2003 Bill was found lacking in these respects and a revised bill was proposed in 2005.

The Plant Protection Bill, 2005, drafted with the assistance of FAO, proposes a new cost recovery mechanism to enable rapid response to epidemics of quarantine importance. It introduces pest risk analysis and strengthens the import and export control of plants, plant products and regulated articles. The Department of Crop Protection is designated as the National Plant Protection Organization (NPPO) and is responsible for the implementation of the act. To this end, the NPPO is in charge of surveillance of growing plants (including areas under cultivation and wild flora) and of plants and plant products in storage or in transport, in order to report the occurrence, outbreak and spread of pests, and to control those pests.

The Minister of Agriculture is authorized to appoint inspectors to enforce the act, from among officers of the NPPO or other competent persons. In addition, the minister may delegate certain functions, by statutory instrument, to any specified competent individual or institution, which includes designation of laboratories and competent scientists.

There are some overlaps in the legislative framework which the bill attempts to address in order to avoid institutional conflict and the resulting inefficiency. For example, Section 12 of the Agricultural Seeds and Plant Act (Chapter 28, 1994) authorizes the National Seed Certification Service (NSCS) to establish phytosanitary standards and practices for crops. The NSCS is further authorized to direct that seeds or plants harbouring pests and diseases be destroyed within a specified period of time and in a specified manner. Similar provisions are in the Cotton Development Act (Chapter 30, 1994), Section 12, with regard to the Minister of Agriculture in consultation with the Cotton Development Organization. On the other hand, under Section 36 of the National Forestry and Tree Planting Act, 2003, the minister responsible for forestry, the National Forestry Authority or a district council is authorized to prescribe the measures to be taken to control or eradicate pests in forests and forest products. To eliminate these potentially overlapping mandates, a clause in the 2005 Plant Protection Bill provides for the primacy of that bill in plant protection matters.
3.6. Viet Nam

The regulatory framework for plant health in Viet Nam is elaborate and comprehensive. The main piece of legislation is the 2001 Ordinance on Plant Protection and Quarantine. The ordinance provides for pest surveillance and control by generally referring to the management of injurious pests, including survey, detection, forecasting and warning of pest occurrence, development, distribution and damage. Government Decree No. 02/2007/ND-CP on Plant Quarantine requires the Ministry of Agriculture and Rural Development to develop a list of regulated pests, and refers to pest risk analysis as the basis for elaborating that list. Notably, the decree specifies the rights and duties of plant owners which include: (a) the right to be informed on pest status and assisted with pest control by the competent governmental bodies; (b) the duty to apply appropriate pest control measures as recommended by competent governmental bodies in order to contain a pest; and (c) the duty to report any pest of economic importance to competent governmental bodies.

The ordinance sets forth the mandate to designate areas in which an outbreak of a pest of economic importance occurs. The mandate lies with the Chairman of the People's Committee or the Ministry of Agriculture and Rural Development depending on the location of the outbreak. The ordinance states that when quarantine or alien pests are detected, the competent state bodies shall order appropriate measures to delimit and eradicate such pests and require the owners of regulated articles to apply those measures immediately.

Provisions on import controls, including issuance of import permits, inspection and treatment of consignments at points of entry and post-entry restrictions, are provided for in the ordinance and in the decree. In addition, the Ministry of Agriculture and Rural Development Decision No. 16/2004/BNN-BVTV sets forth, among other things, model documents for: (a) the import permit; (b) the application for phytosanitary inspection; (c) the declaration form at the point of entry; (d) the record of inspection for consignments and other regulated articles; and (e) the authorization to import.

Phytosanitary certification for export is mentioned in the ordinance while procedures for export inspections are regulated in the decree. With regard to the powers of quarantine officers, article 6 of the same decree empowers quarantine officers to enter any place where regulated
articles are found. Offences and penalties are regulated in subsidiary legislation, namely Decree No. 26/2003/ND-CP.

IV. ANIMAL HEALTH

4.1. Ethiopia

The Animal Disease Control Proclamation No. 267/2002 regulates the prevention and control of animal diseases. The proclamation tasks the Ministry of Agriculture and Rural Development with import controls on animals and animal products as well as animal movement restrictions. Import requirements are set forth, including import permits and inspections. The ministry is obliged to establish an emergency preparedness and epidemic surveillance system to contain the spread of animal diseases and prevent the introduction of exotic diseases into the country. The ministry is authorized to declare animal quarantine in areas infected by animal diseases of economic relevance and to order zoosanitary measures. With regard to exports, the ministry is responsible for international veterinary certification and the establishment of disease-free areas. The proclamation also sets out export conditions and procedures that exporters shall follow.

A set of regulatory instruments are in place for meat production and inspections. The Meat Inspection Proclamation No. 274/1970 confers a mandate on the ministry to control and regulate the production, processing and handling of livestock products. The Meat Inspection Amendment Proclamation No. 81/1976 mandates the ministry to issue regulations and establish criteria for livestock production for human consumption, including classification of products and inspection of processing facilities. The Meat Inspection Regulations No. 428/1970 lay down the requirements for setting up abattoirs and commercial establishments dealing with slaughtering, preparation and processing of livestock products for export from or import into Ethiopia.

As of 2003, the Government of Ethiopia has designed an export development strategy which pays particular attention to the promotion of meat and other livestock products. The government is building capacity to comply with international standards, particularly those emanating from the OIE. Within this context, a series of draft regulatory instruments are under development. These drafts incorporate Biosecurity concepts into the new legal framework, such as by streamlining import and export procedures and pooling resources to conduct risk assessment.
4.2. Ghana

In Ghana, the Disease of Animals Act, 1961, gives the Ministry of Food and Agriculture the power to adopt measures to curb the outbreak of animal diseases. The powers which may be exercised by veterinary officers of the Directorate of Veterinary Services under the act, especially in the event of an outbreak, are aimed at the control and avoidance of the spread of animal diseases. No reference is made to international standards. Given Ghana’s membership in the OIE and its international obligations under the SPS Agreement, there is the need to refer to and incorporate the OIE standards, which are the international norms and benchmarks for animal health.

4.3. India

In India, the Livestock Importation Act, 1989, regulates the import of livestock and livestock products which may be affected by "infections" or "contagious disorders". These may be specified by the central government by notification in the gazette. Section 2(d) describes "livestock products" as including meat and meat products of all kinds, milk and milk products, embryos, ova and semen as well as any other animal product specified by the central government.

The Livestock Act empowers customs officials to carry out animal health inspections under Section 11 of the Customs Act, 1962. Section 3-A of the Livestock Act specifically states that the central government may by notification regulate, restrict or prohibit the import of "any livestock product, which may be liable to affect human or animal health".

The act empowers the state governments to make rules on the detention, inspection, disinfection or destruction of imported livestock and on the powers and duties of appointed persons. Based on this delegation of authority, several states have passed animal health legislation.

The Animal Quarantine and Certification Service within the Ministry of Agriculture is responsible for the implementation of the Livestock Act and for export certification. The Ministry of Environment and Forests is entrusted with the task of protection of wildlife health in sanctuaries and wildlife parks in accordance with the Wildlife (Protection) Act, 1972. Each state government has the power to protect the health of animals within its own boundaries and
has been empowered by the state enactments mentioned above to set up quarantine stations and testing for diseases.

In case of outbreaks or epidemics, the central government issues notifications and guidelines to control and monitor the disease, and has in several instances set up ad hoc monitoring committees. The Department of Animal Husbandry and Dairying has the task of monitoring and coordinating the various institutions that are engaged in animal health.

4.4. Kenya

In Kenya, the Animal Diseases Act, 2006, regulates animal health. The Director of the Veterinary Services Department (VSD) under the Ministry of Agriculture (now, under the Ministry of Livestock and Fisheries) appoints inspectors for the purpose of implementing the provisions of the act. The director is empowered under Section 5 to declare any area to be infected by a notifiable disease, and can extend, diminish or alter the borders of an infected area. The director may also declare areas free from notifiable diseases and may prohibit the movement of animals from one area to another.

Under the act, the VSD may regulate or prohibit for a period of time the importation or the exportation of animals. The minister is authorized to make animal health rules. Subsidiary legislation under the act elaborates rules on issuance of permits, tests required and certification for importation and movement of animals. It also deals with infected areas and prevention of the spread of disease. Rules have been promulgated under the act dealing specifically with foot-and-mouth disease as well as rinderpest and pig diseases. Those rules however do not directly refer to OIE benchmarks.

4.5. Uganda

The Animal Diseases Act (Chapter 38, 1964) is the main piece of legislation governing animal health. The act defines the animal species and lists the diseases to which it applies. Among other zoosanitary measures, the act provides that animal owners should notify a veterinary officer or administrative officer of any disease outbreak. Once he or she has ascertained the existence and nature of the disease, the veterinary officer must report the matter to the Commissioner of Livestock and Entomology and notify other animal owners in the area. The Cattle Traders Act (Chapter 43, 1964) subjects cattle trading to a licensing regime that is managed by veterinary officers.
4.6. Viet Nam

Legal provisions on the protection of animal health and life are found in several different laws, namely, the Fisheries Law of 2003, the Ordinance on Veterinary Controls of 2004 and the Ordinance on Livestock Breeds of 2004.

The Fisheries Law regulates activities related to aquatic animal and aquatic animal products such as breeding, processing, import and export. Activities that cause adverse effects to aquatic animal breeds are generally prohibited. The law establishes a list of aquatic animal species for which aquaculture is prohibited as well as list of chemicals that are banned in aquaculture. The law also envisages a series of measures that must be taken to protect the living environment for aquatic species as well as to preserve and develop rare aquatic species. The harvesting of rare species requires a permit from Ministry of Fisheries or provincial People’s Committees, as do related activities.

Article 35 of the law states that the Ministry of Fisheries is responsible for developing and implementing: (1) standards for feed used in aquaculture; (2) zoosanitary measures in aquaculture; and (3) a list of banned chemicals in aquaculture.

The Ordinance on Veterinary Controls states that the Ministry of Agriculture and Rural Development and the Ministry of Fisheries are responsible for the prevention of animal diseases as well as the quarantine and treatment of infected animals. Article 23 of the ordinance provides that all animals and products of animal origin, when being transported out of districts, must be quarantined at departure. Articles 28 and 29 of the ordinance have regulations on quarantine for imported and exported animals and animal products. Article 26 establishes requirements for domestic transportation of animals and products of animal origin.

The law assigns the responsibility for the management of veterinary drugs and biological products, including microorganisms, to the Ministry of Agriculture and Rural Development, the Ministry of Fisheries and the People’s Committees.

The Ordinance on Livestock Breeds has some provisions related to animal health and life. It generally prohibits activities that may harm safe animal breeding and regulates some zoosanitary aspects of animal breeding, multiplication and trading. Article 9 of the ordinance prohibits the export of
livestock species of genetic value. The Ministry of Agriculture and Rural Development is responsible for the management of agricultural livestock breeds while the Ministry of Fisheries is responsible for aquatic livestock breeds.

V. INVASIVE ANIMAL SPECIES

5.1. Ethiopia

The 1997 Environmental Policy of Ethiopia calls for action to restrict exotic species, including some potentially invasive plants, from biodiversity hotspot areas. Although the country does not have a stand-alone policy or specific legislation on invasive alien species (IAS), the policy provisions can serve as a basis for future action using the existing legislation on plant and animal health.

5.2. Ghana

In Ghana, the prevention of the introduction, control or eradication of alien species that threaten ecosystems, habitats or endemic species is not the subject of any specific piece of legislation. Nor has Ghana enacted legislation to implement the provisions of the CBD. However, several pieces of legislation along sectoral lines – fisheries, forestry, game and wildlife – exist on the statute books and are used to manage IAS.

5.3. India

The Biological Diversity Act, 2002, contains no provision to deal with IAS, and no mention is made of these species throughout the relevant Indian legislation. In the act, general duties are imposed upon the central government to develop strategies, plans and programmes for the conservation and promotion and sustainable use of biological diversity and to integrate these goals of conservation and sustainability into relevant sectoral, and cross-sectoral plans, programmes and policies.

5.4. Kenya

In Kenya, the Environmental Management and Coordination Act (No. 8 of 1999), establishes a legal and institutional framework for the management of the environment. Section 50 requires the National Environment Management Authority (NEMA) to prohibit and control the introduction of alien species into natural habitats. NEMA is expected to issue guidelines on this function but this is yet to be accomplished.
5.5. Uganda

Under the National Environment Act (Chapter 153, 2000), the National Environment Management Authority (NEMA) is responsible for the review and approval of environmental impact assessments. The list of activities for which the assessment is required includes the introduction of new crops and animals and the introduction of fauna and flora into ecosystems of natural conservation areas. Reportedly, neither NEMA nor the Uganda Wildlife Authority, which implements the Uganda Wildlife Act (Chapter 200, 2000), have an active programme of work on IAS.

5.6. Viet Nam

Like Ghana, Viet Nam does not have legislation systematically addressing IAS but rather manages them through sectoral instruments. Decree No. 58/2002/ND/CP on plant quarantine establishes that the import of all IAS of plant origin is prohibited. In specific cases where the import is for scientific purposes, permission from the Minister of Agriculture and Rural Development may be sought.

Decree No. 109/2003/ND/CP on the Preservation and Sustainable Development of Wetlands bans the introduction of new species which may damage ecosystems or modify the gene pool of animals. Article 6 of the 2003 Aquaculture Law states that the farming of new aquatic animal species requires a permit from the Ministry of Aquaculture.

The 2004 Ordinance on Plant Breeding prohibits the import, breeding and commercialization of IAS that may cause harm to human health, the environment or ecosystems. The 2004 Ordinance on Livestock Breeds contains similar provisions.

VI. BIOSAFETY

6.1. Ethiopia

In an attempt to implement its obligations under the Cartagena Protocol, Ethiopia developed the National Biosafety Framework, which is a set of policy, legal and operational documents that includes a draft Biosafety Proclamation. The draft proclamation establishes procedures of prior notification to and authorization by the Environmental Protection Authority...
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(EPA) for research and development, import, export, transit, handling, contained use, transport, placing on the market, use as a pharmaceutical for humans or animals, use as food, feed or for processing of any genetically modified organism (GMO) or products of a GMO. The proclamation initially envisaged a committee of experts from various regulatory agencies to advise the EPA but this has subsequently been abandoned.

The applicant is required to undertake risk assessment to identify potential risks of GMOs or products derived from GMOs on human and animal health and biological diversity, including socio-economic conditions, cultural norms and the environment in general. A GMO exporter is required to provide evidence of the advance informed agreement of the importing country. The EPA is required to make any application available to the public and technical experts and solicit their comments.

The draft proclamation also requires the identification, labelling and packaging of GMOs or their products subject to any authorization procedure prescribed under the draft proclamation. The EPA is tasked with establishing standards in this regard. The draft proclamation also regulates post-authorization monitoring and inspections.

Criminal sanctions are imposed on offenders who contravene mandatory obligations such as those on notification, risk assessment and compliance with standards.

6.2. Ghana

In Ghana, the draft Biosafety Bill, 2004, is designed to domesticate and implement the Cartagena Protocol. The bill is comprehensive, creating a regulatory regime with accompanying regulations to address permits, financing, monitoring and enforcement, approvals and appeals, public participation and information. Decisions on GMOs are to be based on risk assessment, and the relevant procedures are set out in the fourth schedule to the bill.

The draft bill establishes the National Biosafety Authority (NBA). The functions of the NBA are, among others, to receive, respond to and make decisions on applications filed under the bill and to carry out inspections. Taking cognizance of the fact that biosafety is a multi-institutional activity that cuts across several sectors, the draft bill relies on the expertise of
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existing regulatory agencies by establishing a Technical Advisory Committee (TAC), drawing its membership from those agencies.

The functions of the TAC are to: (a) act as the national advisory committee on matters related to genetic modification of organisms and specifically to carry out risk assessments at the request of the Board of the NBA; and (b) advise the NBA, ministries and appropriate bodies on matters concerning the genetic modification of organisms. Those matters include the introduction of GMOs into the environment, the conduct of specific activities or projects concerning GMOs, the contained use of GMOs, the importation and exportation of GMOs and the preparation of regulations and guidelines. The institutional arrangement proposed by the draft bill (establishment of the NBA and reliance on the existing regulatory agencies) points to a way of resolving the conflicts, gaps and overlaps in the Ghanaian regime on Biosecurity.

6.3. India

In India, several pieces of legislation, either in force or in draft, address GMOs. Food regulations cover labelling and other conditions for sale. The Ministry of Health and Family Welfare has notified draft rules to amend the Prevention of Food Adulteration Rules, 1955, and establish new labelling requirements. The draft rules establish that the manufacture, import, transport, storage, distribution or sale of raw or processed food or any ingredients of food, food additives or any food product that may contain genetically modified material in the country is subject to the approval of and conditions imposed by the Genetic Engineering Approval Committee (GEAC), constituted under the Environment Protection Act, 1986. In cases of import, the importer shall submit documents supporting the approval at the time of import.

The Plant Quarantine Order, 2003, seeks to regulate the import of GMOs of plant origin for the purposes of agricultural research or experimentation. The order requires a permit to be issued by the Director of the National Bureau of Plant Genetic Resources, subject to the approval of GEAC or the Review Committee on Genetic Manipulation (RCGM) within the Department of Biotechnology under the Ministry of Science and Technology.

The order does not cover imports for commercial purposes, which are subject to separate clearances set out in rules that were issued in 1989 by the
Ministry of Environment and Forests under the Environment Protection Act (EPA), 1986. The rules and their accompanying guidelines design a multi-layered decision-making structure involving six different bodies (including GEAC and RCGM) in two different ministries (Ministry of Science and Technology and Ministry of Environment and Forests) over four different phases (pre-research, research, commercial release and post-release).

In 2006, the Ministry of Commerce and Industry notified new regulations for the import of genetically modified products under the Foreign Trade (Development and Regulation) Act, 1992. According to these regulations:

- the import of GMOs/living modified organisms (LMOs) for the purpose of (i) research and development; (ii) food; (iii) feed; (iv) processing in bulk; or (v) for release into the environment is governed by the EPA and the related rules of 1989;
- the import of any raw or processed food or feed or any ingredient of food, food additives or food products that contain genetically modified material and are being used for industrial production, environmental release or field application is permissible only with the approval of the GEAC; and,
- institutes/companies wishing to import GMOs for research and development purposes must submit proposals to the RCGM.

The new regulations further provide that all GMO consignments have to carry a declaration to that effect at the time of import, with provision for penal action under the Foreign Trade (Development and Regulation) Act, 1992, in case of non-compliance.

A liability regime is in place under the EPA. Recently, some policy documents recommended the establishment of a National Biotechnology Regulatory Authority to combine the responsibilities of the several regulatory bodies currently empowered to manage biosafety.

6.4. Kenya

In 1998, prior to Kenya’s ratification of the Cartagena Protocol, non-binding regulations were developed by the National Council for Science and Technology, which was established under the Science and Technology Act (Chapter 250, 1977). To date, they are the main regulatory instruments for GMOs in Kenya and require that the release of GMOs be preceded by the
approval of the National Biosafety Committee (NBC). The relevant regulatory authorities shall undertake risk assessment before making the decision to approve or deny approval for the import. For crops, KEPHIS is the relevant authority, advising the NBC on whether or not to allow imports and on what to do after the assessment.

The draft National Biosafety Bill, 2003, is an attempt to expand the coverage of the draft regulations and give a firm legal basis to biosafety regulation in Kenya. It seeks to align the draft regulations with the Cartagena Protocol. Section 5 of the draft bill establishes the National Biosafety Authority (NBA), whose functions are, among others, to:

(a) receive, respond to and make decisions on applications under the draft bill;
(b) identify national requirements for staff development and capacity building in biosafety; and
(c) keep a record of biotechnology and biosafety activities in Kenya.

The NBA is empowered to approve or reject applications as well as to determine whether or not to carry out risk assessments. The following are activities subjected to the written approval of the NBA:

(a) contained use involving GMOs;
(b) introduction of GMOs into the environment;
(c) importation and placing of GMOs on the market; and
(d) transportation of GMOs through Kenya.

Any decision made by the NBA is subject to review upon the request of a regulatory agency or any applicant where there is new scientific information relating to biosafety of the GMOs or where there has been a change of circumstances. The regulatory agencies are in charge of the following:

(a) monitoring applicants’ activities to ensure that they conform to the law;
(b) informing the NBA of any new information aimed at enhancing the continued safe use of GMOs; and
(c) inspecting and evaluating activities involving GMOs.

The Minister of Science and Technology appoints biosafety inspectors, who have comprehensive enforcement powers under the draft bill.
A challenge for Biosecurity in Kenya is that permits with respect to GMOs have been issued on the basis of these draft regulations and not under legislation in force, since the process of promulgating the draft Biosafety Bill has been protracted. The bill has been under discussion since 2002, while an earlier draft Biosafety Bill of 1999 failed to win approval.

6.5. Uganda

The Uganda National Council for Science and Technology (UNCST), which is a statutory body currently under the supervision of the Minister of Finance, established the National Biosafety Council (NBC) with members from the specialized departments/authorities of the various line ministries. The NBC is tasked with evaluating applications for confined field trials of LMOs and referrals made by any department receiving applications for the import of LMOs (e.g. the Department of Crop Protection, for seeds). The decisions are made within the NBC, while risk assessment is carried out by the competent departments/agencies. The functions and procedures of the NBC are not legislated.

The UNCST has proposed the National Biotechnology and Biosafety Policy, 2006, which notes the inadequacy of the legal framework with respect to regulation of modern biotechnology and related issues. Existing provisions are scattered among several pieces of sectoral legislation and are applied by a number of statutory bodies, each concerned with the fulfilment of its own mandate. Despite Uganda’s ratification of the CBD in 1993 and the Cartagena Protocol in 2001, the provisions of these agreements have not been fully incorporated into domestic legislation nor is there an institution that can address the concerns of both these international instruments. Under the Uganda Biosafety Bill of 2005, the UNCST is proposed as the competent authority for biosafety, with the ministry responsible for the environment as the national focal point to provide coordinated communication on behalf of all relevant ministries, departments and agencies. The provisions of the bill are under extensive and still internal revision by the sectoral institutions participating in the NBC.

6.6. Viet Nam

Biosafety is touched upon in some laws of Viet Nam, such as the 2005 Law on Environment Protection, Decree No. 109/2003/ND-CP on the Preservation and Development of Wetlands, the 2003 Ordinance on Hygiene and Food...
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Safety and the 2004 Ordinance on Livestock Breeds. These instruments generally intend to apply the existing legislative framework for conventional processes and products to biotechnology, GMOs and GMO products.

Decision No. 178/1999/QD-TTg on the labelling of domestic and import-export goods requires GMOs to be labelled as such.

In 2005, Regulations on the Management of Biosafety were promulgated by Decision No. 212/2005/QD-TTg under the 2005 Law on Environment Protection. According to the regulations, the Ministry of Natural Resources and Environment is the primary authority responsible for biosafety management at the state level. Other ministries such as the Ministry of Fisheries, the Ministry of Agriculture and Rural Development and the Ministry of Health are responsible for biosafety at the ministerial level. According to the regulations, enterprises shall obtain a biosafety certification upon adoption of risk management measures. For all other aspects of biosafety (e.g. authorization to import, risk assessment), the regulations lack detailed provisions.

A draft Law on Biodiversity is under development which contains a chapter on biosafety. Since the draft law is still subject to extensive revision, its provisions have an uncertain future.

VII. INSTITUTIONAL SET-UP

7.1. Ethiopia

In general terms, the current institutional framework in the five examined areas lacks the necessary coordination to implement a Biosecurity approach. Food safety matters fall within the mandate of several authorities. The leading government institutions responsible for food safety include the Ministry of Health (MOH), which implements the 2002 Public Health Proclamation, the Ministry of Agriculture and Rural Development (MOARD), the Quality Standards Authority of Ethiopia (QSAE), the Ministry of Trade and Industry and the Ethiopian Manufacturing Industries Association. Since 2002, these bodies have established a Technical Committee that implements international standards on food safety systems.

A number of institutions are assigned, under the proclamations establishing them, to undertake food safety inspections in the country. Some of these
institutions are the MOH, the QSAE, the Ethiopia Health and Nutrition Research Institute and the Customs Authority. To strengthen collaboration, the Technical Committee has established the National Food Safety Council, a consultative body of members drawn from regulatory bodies, research institutes, industry, consumers and higher learning institutes involved in food safety.

Reportedly, some conflicts exist between the MOH and the QSAE. While the MOH sees the role of the QSAE as merely procedural in the development and approval of standards, the QSAE claims not only a technical mandate for standards as a regulatory body but also an implementing role with respect to inspections and enforcement.

Animal health activities are carried out by the Animal and Fisheries Resources Development and Regulatory Department (AFRDRD) within the MOARD. Within the AFRDRD, the Veterinary Services Team is responsible for maintaining the safety of food products of animal origin.

The Crop Protection Department of the MOARD is responsible for plant health matters. It has a body of inspectors that are assigned to quarantine stations and border posts. Lack of capacity is accompanied by institutional conflicts. At present, there is a conflict between the Crop Protection Department and the Institute of Biodiversity Conservation (IBC). The latter is vested by the Proclamation on Access to Genetic Resources and Community Knowledge No. 482/2006 with the responsibility of granting access to genetic resources under certain conditions. Although the subject of the proclamation is access to genetic resources and not plant health, every export of plants or plant products is interpreted as constituting access granted on the germplasm embodied in those plants or plant products, thus requiring the consent of the IBC. This often leads to a conflict between the operating procedures of the two entities. A memorandum of understanding or a legislative instrument could resolve the conflict.

As seen above, the 2006 draft Biosafety Proclamation has the Environmental Protection Authority as the implementing institution.

7.2. Ghana

In Ghana, Biosecurity issues are not the responsibility of one ministry, department or agency of state. Instead, several bodies have responsibility for, or are engaged in, activities in this area. These include:
(a) the Ministry of Finance;
(b) the Ministry of Food and Agriculture (MOFA);
(c) the Ministry of Health;
(d) the Ministry of Local Government, Rural Development and Environment; and
(e) the Ministry of Trade, Industry, Private Sector and Presidential Special Initiatives.

Under each ministry, different departments, agencies and institutions operate. Among them:

(a) the Food and Drugs Board (FDB) and the Ghana Standards Board (GSB) are responsible for food safety;
(b) the Directorate of Veterinary Services (DVS) is responsible for animal health;
(c) the Plant Protection and Regulatory Services Directorate (PPRSD) is responsible for plant health;
(d) the Environmental Protection Agency (EPA) is responsible for environmental matters;
(e) the Customs and Excise Preventive Services (CEPS) is responsible for ports and borders in collaboration with the other agencies;
(f) district, municipal and metropolitan assemblies collaborate with the regulatory agencies at the local level especially in monitoring and enforcement at markets; and
(g) the Cocoa Research Institute operates independently from any other agency for quality control and export purposes under the Ministry of Finance.

Some of these departments/agencies, such as the EPA and CEPS, are statutory bodies, and their functions or mandates are provided for in the enactments that established them.

Three institutions are responsible for implementing Ghana’s obligations under the SPS Agreement: the PPRSD is the mandated NPPO; the FDB and the GSB are the implementing agencies for Codex and the WTO TBT Agreement, respectively; and the DVS is the OIE contact point.

As highlighted earlier, the current institutional arrangements for Biosecurity in Ghana are bedevilled by gaps, overlaps and conflicts in the mandates of the
various institutions involved in Biosecurity activities. The border phytosanitary controls are within the mandate of the PPRSD but operative coordination with CEPS is problematic and not legislated. The meat inspection function is a source of conflict/overlap between the DVS, the FDB and the public health officers of the metropolitan and district assemblies. The relationship between the FDB and the GSB is another source of conflict.

The reported tensions between the FDB and the DVS arise from the draft Meat Inspection Bill. Under Part I of the draft bill of 2004, the DVS is designated as the authority responsible for the control of meat hygiene, for all decisions relating to human health and animal health at admission of slaughter animals to the abattoir and for ante-mortem and post-mortem inspections. Because the DVS does not have sufficient staffing, it cannot perform this function efficiently and effectively without the assistance or collaboration of officials from other ministries, departments and agencies.

Another competence conflict exists between the FDB and the GSB on food safety. The 2006 draft Food and Drugs Bill attempts, among other things, to deprive the GSB inspectors of any role with respect to food quality inspection (through a definition of "food" that explicitly offsets the ambit of the Standards Act). However, the 2006 draft Standards Bill retains the food inspection role of GSB inspectors.

With regard to plant health, implementation of the legislation is the responsibility of the PPRSD. The plant health officers carry out phytosanitary inspections at all border points including ports and the international airport. Officers of CEPS, also stationed at the borders, are obligated to notify the PPRSD of any inspected plant materials in imported shipments or baggage. However, in many cases shipments of plant materials are released into the country without PPRSD having inspected the shipments or even having been informed that the shipments have arrived.

As for biosafety, the Board of the National Biosafety Authority has representation drawn from both the public and private sectors. Draft biosafety legislation provides for sectoral representation on the Technical Advisory Committee (TAC), which would help ensure the much-needed coordination and cross-sectoral management required in such a multi-institutional endeavour as Biosecurity. It will also help address the overlaps, conflicts and gaps in the mandates of the various regulatory agencies that will operate within the TAC.
7.3. India

The *biosecurity* legal and institutional framework in India is elaborate. A plethora of laws dealing with *biosecurity* have been enacted with differing objectives and public concerns in mind. Along the same lines, the different legislative instruments are implemented by different institutions. Though disparate, the existing legislative instruments still serve an essential function in specifically addressing the sectoral concerns.

The following table provides a comprehensive overview of the institutional set-up of *biosecurity* in India.

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<tr>
<th>PARENT MINISTRY</th>
<th>INSTITUTION</th>
<th>FUNCTIONS</th>
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<tr>
<td>1. FOOD SAFETY</td>
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<tr>
<td>1. Ministry of Agriculture</td>
<td>• Dept. of Agriculture &amp; Cooperation</td>
<td>• Standardization, grading &amp; quality control of agricultural &amp; allied produce;</td>
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<td></td>
<td>• Directorate of Marketing &amp; Inspection</td>
<td>• Administration of Meat Food Products Order.</td>
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<tr>
<td>2. Ministry of Health &amp; Family Welfare</td>
<td>• Central Committee for Food Standards and its Sub-Committees for Framing of Rules/Standards of Food Articles</td>
<td>• Development of standards on, among other topics: (a) labelling; b) pesticide residues; c) food additives &amp; contaminants; d) microbiology &amp; hygiene; e) packaging.</td>
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<tr>
<td>3. Ministry of Food Processing Industries</td>
<td></td>
<td>• General competence for food safety.</td>
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<tr>
<td>4. Ministry of Commerce &amp; Industry</td>
<td>• Dept. of Commerce Agriculture &amp; Processed Food Products Export Development Authority</td>
<td>• Promotion of HACCP and hygiene in the respective food sectors;</td>
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<td></td>
<td>• Cashew Export Promotion Council Coffee Board</td>
<td>• Audit and certification of the HACCP system through accredited certified bodies;</td>
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<td></td>
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<td>• EIC is the official government inspection body for certifying food products for export. Its</td>
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<td>PARENT MINISTRY</td>
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|                | • Directorate General of Foreign Trade  
|                | • Export Inspection Council (EIC)  
|                | • Marine Produce Export Development Agency  
|                | • Spices Board  
|                | • Tea Board  | certificate covers good manufacturing practices and HACCP, a combination of product specifications and requirements for manufacture, transport and shipping. |
| 5. Ministry of Civil Supplies, Consumer Affairs, Food & Public Distribution | • Dept. of Food & Public Distribution  
| | • Dept. of Consumer Affairs  
| | • Bureau of Indian Standards (BIS)  | • Certification;  
| | | • Licensing of manufacturers. |

II. ANIMAL HEALTH


2. Ministry of Agriculture | • Animal Quarantine & Certification Service  
| | • Dept. of Animal Husbandry & Dairying  | • Regulation, restriction and prohibition of the import of livestock which may affect human or animal health;  
| | | • Coordination with state authorities. |

III. PLANT HEALTH

1. Ministry of Agriculture | • Dept. of Agriculture & Cooperation  | • Regulation of domestic and international movement of pests, insects and fungi which might threaten agriculture;  
| | | • Regulation of seed quality. |

2. Ministry of Environment & Forests | • Conservation & Survey Division  | • Regulation of access to and conservation of the biological resources of the country. |
Food safety is under the umbrella of the Ministry of Health and Family Welfare, while the Directorate General of Health Services is the Codex Contact Point and works in collaboration with the Ministry of Food Processing Industries. As mentioned above, the official standard-setting authority is the BIS, which also has quality certification functions. Export certification is ensured by the EIC.

The Food Safety and Standards Act, 2006, establishes the Food Safety and Standards Authority of India (FSSAI), which is assisted by a central advisory committee, a scientific committee and several scientific panels. The Commissioner of Food Safety of each state enforces the standards through food safety officers. The FSSAI is mandated to lay down science-based standards for food articles, and seeks to regulate their manufacture, import, storage, distribution and sale to ensure availability of safe and wholesome food for human consumption.

The Commissioner of Food Safety of each state appoints a designated officer for a specific district whose duties include issuing or cancelling licences, prohibiting sale of food articles that violate specified standards, receiving reports and samples of food articles from food safety officers and having them analyzed.

With regard to animal health, the main authorities are the Department of Animal Husbandry and Dairying and the Animal Quarantine and Certification Service in the Ministry of Agriculture. However, wildlife management in protected areas falls under the Ministry of Environment and Forests. The need for a more effective centralized authority to monitor and coordinate the various activities of the state authorities is clear.
Plant health is under the Department of Agriculture and Cooperation in the Ministry of Agriculture. The Ministry of Environment and Forests has biodiversity-related functions that complete the Biosecurity framework.

Although there are a variety of institutions, there is a general appreciation of the sectoral work of the relevant institutions while the concept of a single agency is viewed with caution. A gradual upgrading of the legal framework, tailoring the mandates of existing institutions to carry out their Biosecurity-related tasks, may be the better approach for India.

7.4. Kenya

The institutional basis for Biosecurity is shared among different ministries and institutions in Kenya. These include:

(a) the Ministry of Agriculture, under which the Kenya Plant Health Inspectorate Service (KEPHIS) operates;
(b) the Ministry of Livestock and Fisheries, under which the Veterinary Services Department (VSD) operates;
(c) the Ministry of Environment, under which the National Environment Management Authority (NEMA) is established;
(d) the Ministry of Trade and Industry, which oversees the Kenya Bureau of Standards (KEBS);
(e) the Ministry of Education, Science and Technology, under which the National Council for Science and Technology (NCST) and the National Biosafety Committee (NBC) were created; and
(f) the Ministry of Health and its Central Board of Health.

In accordance with its technical mandate, each institution is responsible for Kenya’s international obligations under the SPS Agreement.

To bring together the institutions responsible for different regulatory functions, a number of inter-ministerial coordinating committees have been established. The National Committee on WTO, for instance, gathers KEBS, KEPHIS and NEMA. The committee is established under the Ministry of Trade and Industry, and the ministries sitting on the committee act as focal points for sub-committees handling relevant WTO issues falling within their mandates. Another example is the Kenya Standing Committee on Imports and Exports.
The establishment of KEPhIS in 1996 has led to greater coordination of the phytosanitary aspects of Biosecurity. This has however been hampered by delay in amending the relevant laws to legitimize the role of KEPhIS. It is a matter of particular concern that a proposed bill to institutionalize KEPhIS, drafted with the assistance of FAO, is yet to be promulgated and continues to be debated. The central role of KEPhIS in the Biosecurity framework in Kenya suffers from the uncertainty of its legal basis. In fact, having been established by ministerial order and not by parliamentary-level legislation, it could, at least in theory, be dissolved or its role changed at any time by a new ministerial order.

Furthermore, there has been delay in amending the relevant sectoral laws to be implemented by KEPhIS. The Plant Protection Act (Chapter 324, 1962, as amended), the Suppression of Noxious Weeds Act (Chapter 325, 1986) and the Seeds and Plant Varieties Act (Chapter 326, 1972, as amended) all need to be amended to synchronize their provisions with the role of KEPhIS. Currently, there is room for conflict between the Ministry of Agriculture officers and KEPhIS officers in the performance of their duties. However, a positive trend is the involvement of officers from line regulatory institutions in related regulatory bodies such as the involvement of KEPhIS, the VSD and KEBS in NEMA and the NBC. This has assisted in coordinating various Biosecurity functions.

With regard to food safety, the Food, Drugs and Chemical Substances Act (Chapter 254, 1970) is implemented by the Ministry of Agriculture, which is also responsible for veterinary services. Under the Meat Control Act (Chapter 356, 1973), some of the inspectors’ activities require consultation between the Ministry of Agriculture and the Ministry of Health. However, the limited human resources available in the ministries make those consultations rare and formal. Moreover, because the Meat Control Act deals with meat for export, the ministry responsible for trade and industry is also a relevant player in the implementation of the act, especially for export certification.

Recently, the need to streamline the food safety aspects of Biosecurity has been recognized and a Food Safety Committee set up, with the Agriculture Secretary as chair. The committee, launched on 4 May 2007, is the focal point for all food safety issues and draws its membership from the Ministries of Agriculture, Health and Trade, the Kenya Medical Research Institute, KEBS and the Kenya Agriculture Research Institute.
Food standards are established by KEBS, which was established under the Standards Act (Chapter 496, 1974). KEBS works closely with the main public bodies in the development and implementation of health standards on animals and animal products, plants and plant products and food safety. The main public bodies are KEPHIS, the VSD and the Ministry of Health. KEBS is the contact point for Codex and the International Organization for Standardization (ISO).

As for plant health, the regulatory agents under the Seeds and Plant Varieties Act include the Minister of Agriculture, seed analysts and KEPHIS. Ordinarily, KEPHIS seed inspectors perform tasks under the Plant Protection Act and the Suppression of Noxious Weeds Act. Under the latter act, local authorities are empowered to make by-laws regarding the eradication of any noxious weed from land within their areas, appointing inspectors and compelling owners or occupiers of land to eradicate any such weed from their land.

With regard to animal health, the Animal Diseases Act (Chapter 364, 1972, as amended) vests the VSD with the power to appoint inspectors. Reportedly, the mandate of KEPHIS includes the enforcement of standards for good husbandry and the control of animal diseases, although these powers do not appear in any legislative instrument.

In general, inspections are a problem arising mainly from scarcity of resources, which impedes the effective discharge of duties entrusted to the officers. For instance, while KEPHIS is required to provide border control with respect to plant materials (a function previously performed by airport staff of the Ministry of Agriculture), there are not enough inspectors to cover all entry points. Moreover, there are no inspectors of the VSD to inspect meat and meat products at the points of entry. Proposals to establish a single agency to encompass both KEPHIS and VSD have been made over the years but have not been implemented.

Environmental legislation is the most advanced in terms of institutional provisions. To ensure compliance, the Environmental Management and Coordination Act (No. 8 of 1999) establishes an elaborate institutional framework. Under the act, the National Environment Council is responsible for formulating policy on matters relating to environmental management in Kenya. It sets national goals and objectives and determines policies and priorities for the protection of the environment. It also promotes
cooperation among public departments, local authorities, the private sector, non-governmental organizations and other organizations engaged in environmental protection programmes. NEMA is the principal government institution responsible for the implementation of all policies relating to the environment. It is also responsible for dealing with environmental impact assessments, including from LMOs and IAS.

Similar to the situation in Ghana, the most detailed institutional provisions in Kenyan legislation appear in the draft Biosafety Regulations of 1998 and the draft National Biosafety Bill of 2003. The draft bill establishes the National Biosafety Authority, managed by a board drawing from the main agencies dealing with biosafety as well as other scientific experts and a consumer representative.

7.5. Uganda

As in the other countries studied, Biosecurity issues in Uganda are the responsibility of different ministries, state agencies or departments. The pieces of legislation which address food safety, plant and animal health are sectoral in nature, and different departments or regulatory agencies are responsible for their implementation.

The Ministry of Health (MOH) is the implementing authority of the Food and Drugs Act (FDA) (Chapter 278, 1964), but it is commonly recognized that its human and financial resources are extremely limited and food safety is not prioritized. Funding is sporadic, meaning that the MOH generally has a reactive approach to food safety issues. In practice, the MOH has been working with other agencies, on a case-by-case basis, in emergency situations (e.g. the European Union ban on fish from Uganda where MOH worked with the Ministry of Agriculture, Animal Industry and Fisheries (MAAIF) and the Uganda National Bureau of Standards (UNBS) to resolve the issues). A very advanced food chain approach is only implemented for fisheries products thanks to cooperation between the Fisheries Department and UNBS.

The Food Hygiene Advisory Committee, established under the FDA, and the minister, in conjunction with the local governments and authorized persons, are responsible for the prevention of the adulteration of food. UNBS, as the national standards body, sets and enforces standards, in some instances adopting standards from other jurisdictions or from international agencies for application in Uganda. Inspection of food imports is mostly done by UNBS agents at entry points.
With regard to animal health, the veterinary services division of the MAAIF is the main authority and the OIE contact point, while the fisheries arm of MAAIF handles aquatic life issues.

The Department of Crop Protection is responsible for phytosanitary and plant protection matters and will be designated the national plant protection organization once the Plant Protection Bill is enacted. The department also acts as the SPS Enquiry Point.

The National Environment Management Authority (NEMA) has a coordinating, monitoring and supervisory role and is the national focal point for the CBD and related instruments. The Uganda Wildlife Authority is the principal body responsible for wildlife management in Uganda, while the National Forestry Authority has similar responsibility with respect to forests. The Customs Department of the Uganda Revenue Authority, established in 1991, plays a crucial role in ensuring the legitimacy of imports and exports of regulated materials.

The Uganda National Council for Science and Technology (UNCST) is responsible for setting policy in all fields of science and technology and acts as the national focal point for the Cartagena Protocol. It is the competent authority for regulation and access to genetic resources, and is proposed as the competent authority to supervise and regulate the implementation of the draft policy on biotechnology and biosafety. Furthermore, it is proposed as the competent authority for biosafety under the draft Biosafety Bill of 2005, along with the ministry responsible for the environment. Notably, through its National Biosafety Council, UNCST is already a forum where sectoral authorities converge to deal with cross-cutting issues such as biosafety. This role suggests that UNCST may be the appropriate forum to start discussions on the implementation of a Biosecurity approach.

Each of the agencies mentioned above has inspectors in charge of ensuring compliance with the provisions of the applicable sectoral law. Certain laws also provide for delegation, enabling collaboration between departments and agencies. In appointing environment inspectors, NEMA is authorized to gazette persons employed as inspectors in other departments. NEMA itself, however, lacks the human resources to participate in activities of other ministries at the technical level.
Similarly, the phytosanitary service cooperates with the Customs Department in undertaking phytosanitary inspection at the various entry and exit points. Inspection of meat is undertaken by veterinary surgeons, medical officers or health inspectors authorized by the relevant minister or the local authority. However, most inspectors are only trained in their particular field of expertise and do not have sufficient capacity to effectively undertake inspection by delegation.

Finally it is worth noting that, although UNBS is the national standards body, other government units have the authority in law to set and enforce standards. For example, NEMA sets environmental standards, while the Directorate of Water Development sets water and water-related standards under the Water Act (Chapter 152, 1995).

7.6. Viet Nam

The institutional set-up of Viet Nam is characterized by a somewhat rigid division of responsibilities among the line ministries for each of the **Biosecurity** areas. The Ministry of Science and Technology is responsible for promulgating standards in all areas of **Biosecurity**, and is also the Codex Contact Point. The Ministry of Health is responsible for formulating and promulgating strategies and policies on food hygiene and safety as well as taking an oversight role in the prevention of food poisoning. It is in charge of the development of regulations on food hygiene and safety and their enforcement. In emergencies, it coordinates with the People’s Committees at all levels as well as the concerned ministries to establish control measures.

The Ministry of Science and Technology issued Decision No. 25/2004/QD-BKHCN promulgating the statute of the Viet Nam Codex Commission. Pursuant to this decision, the Vietnam Food Standardization Committee acts as the Viet Nam Codex Commission, chaired by the Ministry of Science and Technology. The Directorate for Standards and Quality under the Ministry of Science and Technology is part of the Viet Nam Codex Commission and acts as the Codex Contact Point.

The Ministry of Agriculture and Rural Development (MOARD) oversees the production processes of food and the control of hygiene of imported food of animal origin. The Ministry of Fisheries (MOF) is responsible for aquatic products for domestic consumption and aquatic food products which are exported or temporarily imported for re-export. MOF, in coordination with other relevant ministries, is responsible for inspecting enterprises and
monitoring compliance with regulations on the quality and safety of imported and exported fish products.

MOARD establishes and implements quarantine measures for animals and plants, while MOF is responsible for sanitary measures on aquatic animals. In this regard, a resolution of the National Assembly adopted in August 2007 provides for the merger of the two ministries.

With respect to IAS and biosafety, the Viet Nam Environment Protection Agency takes the lead under the Ministry of Natural Resources and Environment, which hosts the focal point for the CBD and the Cartagena Protocol.

Viet Nam also has an inter-ministerial working group that coordinates SPS activities. Decision No. 99/2005/QD-TTg of the Prime Minister established the Viet Nam Sanitary and Phytosanitary Notification Authority and Enquiry Point, which serves as enquiry point and notification authority under the SPS Agreement.

VIII. CONCLUSION

This chapter has presented only a snapshot of the main areas of Biosecurity, to show the analysis as it has been carried out in the six pilot countries. The comprehensive reviews of the Biosecurity frameworks in each of the six countries are presented in Chapters 5–10.

The gap analyses presented here clearly demonstrate the methodology required to assess national frameworks for Biosecurity. The review requires an analysis of the legislation covering the main sectors of Biosecurity – food safety, plant health, animal health, invasive animal species and biosafety – assessing whether the legislation follows international standards and whether there are overlaps and gaps. Next the investigation must turn to the institutional set-up, examining which institutions are empowered to carry out Biosecurity functions and where there are any weaknesses, such as duplications of responsibilities or unclear mandates.

The analysis developed here is refined in the next chapter, which sets out in detail the methodology and its constituent steps. Application of this methodology should lead to a more comprehensive understanding of the national legal and institutional frameworks for Biosecurity and identification of any corrective action needed.
LEGAL METHODOLOGY

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I. OVERVIEW

As can be seen from the review of the pilot case studies, a Biosecurity legal assessment consists of the following two steps:

1. an analysis of the legal framework covering the subject areas of Biosecurity;
2. a review of the mandates and functions of the various institutions responsible for Biosecurity controls.

Based on that analysis, policymakers will then need to consider the feasibility of taking three additional steps:

3. creating new agencies or establishing coordination mechanisms to implement a government-wide Biosecurity approach;
4. elaborating a legislative strategy to pursue the Biosecurity approach; and
5. implementing the strategy through new or amended legislation.

The next sections expand on the methodology to be applied in each of the five steps.

II. ANALYSE SECTORAL LAWS

The first step in the Biosecurity methodology will be an analysis of the existing legislative framework for Biosecurity. The logical division of subject areas for review is the following:

(a) food safety;
(b) plant health;
(c) animal health;
(d) invasive alien species; and
(e) biosafety.

Although every effort should be made to identify the main purpose of each statute so as to fit it into one of the subject areas above, the distinction between certain areas may be blurred, such as between animal health and food safety in cases of legislation covering animal products such as meat. The same can occur with plant health legislation and rules governing the export of fresh agricultural produce. Other laws, such as laws establishing a
national standard-setting authority (e.g. the Quality Standards Authority of Ethiopia, the Bureau of Indian Standards), may also need to be considered carefully as they may not fall neatly into one category. In fact, standard-setting is a typically cross-cutting mandate that intersects with the responsibilities of Biosecurity institutions, most frequently food safety, animal health and plant health authorities.

In each of the sectoral areas of Biosecurity, government officials interested in carrying out a review of the national legal framework should first collect and then analyse the various pieces of relevant legislation. The review of existing legislation should cover both parliamentary-level and subsidiary legislation. Typically, for food safety, animal health and plant health, there is one main parliamentary-level legislative instrument setting forth the general discipline (for instance, the Food and Drugs Law in Ghana or the Plant Protection Act in Uganda). This legislation often contains very brief provisions assigning the mandate for certain activities (for instance, quarantine inspections at border posts or certifications for export) to an authority.

The basic law is (or should be) accompanied by subsidiary legislation which specifies exactly how the assigned mandate is to be executed and provides some operational details (such as time frames for inspections, duties to notify competent authorities of arrival of consignments, model forms and certificates). It is necessary to look at those details, if provided for in subsidiary instruments such as ministerial regulations and orders, in order to carry out a comprehensive review of the strengths and pitfalls of the legislation in that sector. For instance, if a law has provisions on import requirements to be published in regulations but those regulations are either not present or outdated, the law may have little or no effect.

With regard to IAS, it is uncommon to find a single piece of legislation covering the subject. IAS can have different origins (plant or animal) and can be categorized in several ways. For instance, some IAS may qualify as plant pests and be regulated under phytosanitary legislation while others may not be regulated at all.

In biosafety, it is frequent to find biosafety laws, sometimes still in draft form, that are patterned after the Cartagena Protocol (see those examined in Chapters 8 (Kenya) and 9 (Uganda)). The regulation of living modified organisms may also be scattered in several pieces of legislation (see Chapter 10 (Viet Nam)).
Once the legislation is collected and classified according to its subject matter and status (parliamentary-level or subsidiary), policy-makers will have to evaluate the legislation in each of the Biosecurity areas. This evaluation has two parts. The first evaluates whether the legislation covers all the relevant sub-topics in that sector, as there are certain legislative provisions that must be present in order to build a comprehensive regulatory framework for that Biosecurity sector. The second looks at whether the legislation meets international obligations.

The next section provides a brief overview of the main sub-topics which should be addressed in each of the Biosecurity sectors.

2.1. Substance of sectoral laws

With regard to food legislation, the law should determine what kinds of food it regulates (e.g., foods of animal origin, street foods) and what harmful substances in food (e.g., food additives, pesticides, veterinary drug residues) it covers. The legislation should cover food hygiene by setting out the basic principles and rules to be followed by owners and operators of food establishments during the preparation, processing, manufacturing, handling, packaging, transportation, storage and distribution of food in order to guarantee a safe product fit for human consumption.1 The food legislation should contain substantive provisions in these areas, stating for instance that food businesses shall follow hygiene rules and food manufacturers shall establish trace-back procedures.2 Food legislation should also contain rules applicable to imported and exported food (such as the requirement to seek a permit from the competent authority).

Plant health legislation should be designed so as to guarantee that the government can create or function as an effective administrative and technical structure (the National Plant Protection Organization (NPPO)) for the implementation and enforcement of phytosanitary measures. The legislation should allow the NPPO to take action to control the introduction and spread of certain pests which will be listed in the legislation in accordance with risk analysis. Provisions for the establishment of quarantine areas are essential in order to contain outbreaks of quarantine pests.

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1 J. Vapnek and M. Spreij, Perspectives and Guidelines on Food Legislation, with a New Model Food Law, FAO Legislative Study No. 87, 2005, p. 86.
2 Id. p. 173.
Phytosanitary legislation should also address the many aspects of the import and export of plants and plant products. For imports, the phytosanitary legislation should provide for the establishment of import requirements and phytosanitary measures based on scientific justification, including measures to be taken in emergency situations. With regard to exports, the legislation should provide for phytosanitary certification by the NPPO. Furthermore, the legislation should contain provisions allowing the NPPO to take action for detection, survey, containment and eradication of plant pests within the territory, and should provide for the establishment and maintenance of pest free areas.3

Animal health legislation should cover terrestrial as well as aquatic animals, and should set out a list of diseases and national pathogens based on risk analysis. The law or laws should charge the government with preventing and controlling these diseases and pathogens, through surveillance, monitoring, official control and stamping-out programmes. Other activities to be regulated include emergency action in case of disease and pathogen outbreaks as well as animal identification, traceability and movement. Animal health legislation should also permit the government authorities to establish buffer zones, free zones, zones of low disease prevalence and surveillance zones for zoosanitary purposes.

The legislation should also regulate the import of animals and animal products. The issuance of international veterinary certificates for the export of animals and animal products is another key regulatory area. In response to modern developments on animal welfare, the establishment of standards during the life of an animal as well as during its slaughter and destruction can be addressed in the legislation. Tangential but also important issues which may be regulated in separate laws or regulations are provisions on the import and export of animal feed as well as the manufacture, import, export, use, quality, suitability, packaging, labelling, transport, storage, sale and advertising of veterinary drugs.

The regulation of IAS may be found in several pieces of legislation, such as legislation on plant protection, biodiversity or nature conservation. The legislation will have to address prevention and containment of any introduction or invasion of IAS, covering risk management for those species.

As IAS is a cross-cutting subject, it will be important to ensure that the various pieces of legislation do not cause any overlap or conflict in oversight activities, for instance between plant health and biodiversity authorities.

With regard to biosafety, the legal framework should first define its scope (what activities and organisms are covered) and establish an authorization/licence/permit system for those activities and organisms. The legislation should set out the necessary specifications for the system, such as the information to be provided by the applicants, requirements for applications and time limits for decisions. The legislation may also set out simplified procedures for low-risk categories of living modified organisms (LMOs) as well as requirements for public consultation on permit applications. The legislation will also need to address risk assessment procedures and criteria, risk management conditions which may accompany the permit and post-approval monitoring and review. It should also address cases of unintentional releases of LMOs and emergency measures applicable in such circumstances.

2.2. Conformity with international obligations

The second objective of the legal assessment is to evaluate the conformity of the national legal framework with international dictates. In this exercise, the task will be to assess if and to what extent national legislation enables full compliance with the country’s international obligations. This does not mean that international obligations need to be specifically spelled out in national laws. Rather, the task for national authorities is to develop national laws in light of and in harmony with the international instruments which are applicable at national level. Ultimately, what matters is how national measures are executed in practice and, in the case of the sanitary and phytosanitary measures, how transparently they are established and applied. It is against this criterion and not through the literal transposition of clauses of international agreements into national laws that the domestic legal framework should be evaluated.

A few examples from the pilot case studies show provisions of national legislation that are clearly not in line with international standards. In the area of food safety, Schedule I of Ghana’s Food and Drugs Law, 1992, makes

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reference to the publications of certain international bodies but the list does not include those of the Codex Alimentarius Commission, which is the internationally recognized source of international food standards under the World Trade Organization Agreement on the Application of Sanitary and Sanitary Measures (SPS Agreement). In the area of plant health, Ethiopia’s Council of Ministers Regulation No. 4/1995 does not provide for the establishment of any pest list based on risk analysis, which is the cornerstone of the regulatory set-up of the International Plant Protection Convention (IPPC). With regard to animal health, the legislation of Uganda does not provide for any notification of diseases to the Office international des épidemies (OIE) based on which the OIE could circulate relevant information to the international community.

To carry out a detailed review of the sectoral areas and to evaluate their conformity with international standards, FAO has developed several documents and guidelines which can be of assistance:

- Perspectives and Guidelines on Food Legislation;\(^6\)
- Guidelines for the Revision of National Plant Protection Legislation;\(^7\)
- Institutional and Legal Measures to Combat African Swine Fever;\(^8\)
- Decision-Support Toolbox for Biosafety Implementation.\(^9\)

Other more general assessment tools contain useful guidance, including:

- Phytosanitary Capacity Evaluation Tool;\(^10\)
- Assuring Food Safety and Quality: Strengthening National Food Control Systems.\(^11\)

The websites of the main international organizations and instruments governing Biosecurity sectors will also have information and guidance for countries engaged in a detailed review of their national legislative framework for Biosecurity.

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6 J. Vapnek and M. Spreij, supra note 1.
7 J. Vapnek and D. Manzella, supra note 3.
11 Assuring Food Safety and Quality: Guidelines for Strengthening Food Control Systems, FAO/WHO, 2003. At the time of writing, FAO was also finalizing a Biosecurity Capacity Assessment Tool that is part of a Biosecurity Toolkit.
III. REVIEW INSTITUTIONAL MANDATES

Reviewing the legislation in each Biosecurity sector is only the first step in the assessment of the Biosecurity framework. Governments must look at the overall picture of controls on food safety, plant health and animal health to assess whether there is or can be created an efficient, integrated system to manage biological risks. The pilot case studies clearly bear out that the challenge is to foster coordination among regulatory bodies so as to eliminate gaps, overlaps and conflicts.

In each of the five sectoral areas of Biosecurity, it will be necessary to look closely at the institutions implementing the sectoral laws. In assessing the institutional mandates, there are four main issues to consider:

1. what types of institutions carry out Biosecurity controls;
2. whether the responsibilities of the different institutions are legislated and, if so, at what level (parliamentary-level or subsidiary legislation);\(^{12}\)
3. whether there are gaps or overlaps in the exercise of the functions; and
4. whether institutions correct any overlaps or gaps in the legislation through de facto arrangements.

On the first issue, the Ghana study shows that, whereas the Food and Drugs Law establishes the Food and Drugs Board as a statutory body, the Prevention and Control of Pests and Diseases of Plants Act assigns the mandate for plant health control to the Ministry of Food and Agriculture in general, under which the Plant Protection and Regulatory Services Department (PPRSD) operates as an internal department without any specific legislated powers (see Chapter 6). This has implications not only for the different capabilities of the two institutions in terms of material and human resources, but also for the relative deference that other national authorities (such as customs) and the private sector (such as importers and exporters) accord the two bodies.

\(^{12}\) The legal force of the legal instrument under which a body is established is important first because certain powers can only be provided for in primary legislation as they require parliamentary approval. Second, in principle, legal instruments can only be repealed by the same or higher-level instruments, hence a ministerial decree can easily be replaced by a new ministerial decree, for instance where a new minister is appointed, while a parliamentary-level law can only be amended by parliament.
Similarly, the Veterinary Services Department in Kenya remains a
department of the government while the Kenya Plant Health Inspectorate
Service (KEPHIS) is a state corporation with more autonomy and flexibility
(see Chapter 8). In India, food safety is managed by a statutory body (the
Food Safety and Standards Authority), while the main implementing
authorities for plant and animal health (respectively, the Department of
Agriculture and the Animal Quarantine and Certification Service) are
ministerial departments (see Chapter 7). These may be deliberate choices by
government to emphasize certain sectors over others, but to foster
consistency across Biosecurity subject areas may require a closer look at such
institutional anomalies.

On the second issue (level of legislation), the Kenya study highlights that
some of the responsibilities that KEPHIS exercises at present do not have a
basis in law while others are set out only in subsidiary instruments (see
Chapter 8). The chapter notes that a parliamentary act establishing KEPHIS
as a statutory body and defining its overall mandate is still being debated.

On the third issue (gaps and overlaps), the question is whether certain
regulatory functions fail to be implemented by any institution or, conversely,
whether there are areas where two or more institutions overlap. An example
of the first case is Uganda, where the Food and Drugs Act tasks the Ministry
of Health with food hygiene standard-setting and controls, but it is presently
not implementing these responsibilities (see Chapter 9). For the second type
of problem, the meat inspection dispute between Ghana’s Food and Drugs
Board and the Directorate of Veterinary Services of the Ministry of Food
and Agriculture is exacerbated by conflicting legislative provisions.

In carrying out the assessment of the various Biosecurity-related institutions, it
may be useful to look at the following Biosecurity functions:

- diagnostic services;
- quarantine services;
- surveillance and monitoring;
- emergency action;
- inspection services;
- scientific research and advice;
- enforcement.
The task will be to determine whether the institutions are empowered by the law to carry out all of these functions in each Biosecurity sector. For instance, plant health legislation may impose a duty on customs officials to notify the arrival of plants or plant products to plant health authorities, and they do so routinely, but no such duty is set forth in animal health or food safety legislation and no such notification takes place. This is a clear gap in Biosecurity controls as a whole. The function-based analysis will also assist in devising legislative solutions to improve the integration of Biosecurity controls among the sectoral authorities.

The final step is to look beyond how the legislation creates or empowers the different institutions to any other arrangements in place. In many cases there may be written agreements (such as memoranda of understanding) or even informal arrangements between two or more institutions to correct specific gaps or overlaps in mandate or functions. For instance, although neither Ghana’s Prevention and Control of Pests and Diseases of Plants Act nor any subsidiary legislation provides for consultation with environmental authorities, the PPRSD has taken initiative and regularly involves academic environmental experts in the pest risk assessment of certain plants and plant products (see Chapter 6). A similar example is the Department of Crop Protection in Uganda, which regularly involves environmental experts appointed with the concurrence of the National Environment Management Authority for its pest risk analysis in the absence of any internal memorandum formalizing the arrangement (see Chapter 9). Similarly, the National Biosafety Council of Uganda receives and acts upon applications for the import of LMOs of plant or animal origin referred to it by the Department of Crop Protection and the Veterinary Services Department, without specific authorization under any legal text.

IV. CONSIDER CREATION OF A NEW AGENCY/COORDINATION MECHANISM

Having identified the weaknesses and gaps in the legislative and institutional frameworks, governments will next need to consider possible solutions in order to integrate Biosecurity functions and rationalize the Biosecurity framework. One option is to create a new body at a supra-ministerial level, for example under the Presidency or the Council of Ministers, to implement a Biosecurity approach. Another option is to use existing legal and institutional frameworks while establishing a coordinating mechanism to exercise an oversight role.
The creation of a new body along the lines just mentioned has the merit of according the subject the attention that it deserves at the highest level of government. This should enhance its effectiveness and prevent "inter-ministerial jealousies", since *Biosecurity* would not be under the control of just one ministry. The unified agency would have responsibility for end-to-end *Biosecurity*, overseeing pre-border and border activities, incursions and eradications and pest and disease management. The agency would also coordinate with any authority responsible for biodiversity protection in relation to IAS and to biosafety.

The prominent role of the lead agency, of course, does not mean that it would work alone. From time to time, it may need to delegate tasks to other departments or units where there is specialized knowledge. It should also develop systems to protect wider interests in *Biosecurity* and improve connections among the agricultural, environmental and health sectors.

The creation of a new agency has the advantage of mitigating the problem that in some countries, not all *Biosecurity* sectors are at the same stage of development. For instance, in some sectors there may be critical gaps in baseline knowledge while others may lack diagnostic and treatment tools and equipment. Where centralization into a new agency occurs, capabilities, in terms of material resources and intellectual capital, could be maximized through resource pooling (e.g. at the border posts) instead of having isolated units with minimal staffing and equipment addressing narrow sectoral concerns. Although the idea of creating a new institution may be daunting, it can actually be well suited to developing countries and small states with resource constraints.

Despite the potential advantages, the creation of a new institution to address *Biosecurity* may still not be possible in some national contexts. There may be political resistance, for instance due to the historical separation of certain sanitary and phytosanitary functions or due to particularly powerful ministers resistant to loss of influence. And despite the potential advantages of resource pooling, there is no doubt that creation of a new agency entails heavy financial, logistic and manpower requirements which many countries can ill afford. Thus, another option is establish a coordination mechanism to oversee existing line agencies. Such a mechanism – a board, council or committee – would be the repository of information as well as the mechanism for disseminating information to all relevant actors. The board or council would be given certain powers to oversee the entire framework and
would have the authority to require certain actions by line agencies to ensure effective **Biosecurity** controls.

This solution would of course require line agencies to cede some control to the overall coordinating mechanism, and the regulatory framework may have to be modified to guarantee the much-needed coordination inherent in such a multi-institutional activity as **Biosecurity**. The biosafety area is a good example of successful cooperation in three of the countries examined in this study (namely, Ghana, Kenya and Uganda): each used existing institutions while creating an oversight body for biosafety issues, drawing on the implicated ministries and agencies on the national scene.

As for the actual implementation of **Biosecurity** controls, efficiency in the application of such measures depends on standardized risk assessment and management procedures, which, in turn, rely on science. Identifying the right advice is the key to making good decisions. Scientific input from the best-positioned experts, no matter which institution they serve, must be relied upon, bolstered by public and stakeholder input. Laws can assist in the establishment of science-based criteria for sanitary, phytosanitary and zoonosanitary measures as well as procedures to ensure that those criteria are applied.

### V. DEVISE LEGISLATIVE STRATEGY

Having identified the weaknesses and gaps in the existing legislative and institutional frameworks and having decided upon the institutional set-up to be pursued, the next step is to identify a legislative strategy to implement the necessary changes. Elaborating the strategy will require extensive consultations to verify the feasibility of legislative change. It is important to understand the context in which legislative change will take place, to have a realistic understanding of how open to change decision-makers are in a particular setting.

Several considerations will affect the design of a legal strategy for the particular country. The first is the legal system. Each country has its own history, politics, traditions, legislation, institutions and resources. Any new legislation must be conceived with these factors in view, in order to ensure that the proposed legislation reflects national needs and national circumstances.
Apart from the formal legal system, the role of law in society varies enormously from country to country. In some countries, adopted legislation may be generally effective, while in others it may have little impact, mainly because of lack of resources for implementation and enforcement. The absence of necessary political will to support certain recommendations may also be related to the manner in which the law is perceived by public authorities. Much-needed collaboration among authorities may fall victim to institutional jealousies, turf-defending behaviour and passive resistance of government officials or stakeholder groups. These constituents may feel their interests can be better protected through new sectoral legislation that is promoted autonomously and not as part of a collective and comprehensive Biosecurity approach.

Another important consideration in the design of nationally tailored legislative strategies is the government’s policies and priorities. In every country, a variety of policies, strategies and priorities of national, regional or international provenance affect the development of national legal frameworks. In some situations, governments are obligated to incorporate certain policies in their national legislation, while in others they may do so voluntarily. Hence, undertaking an assessment of national Biosecurity legislation entails understanding the level of commitment that the government has with respect to Biosecurity in the context of other relevant policies.

As happens with laws, policies have varying degrees of importance in different countries and this will be an important consideration in elaborating a legislative strategy. If a decentralization policy or decentralization law is extremely influential, this will affect the design of the legislation in fundamental ways. Thus, in any new legislative framework for Biosecurity control, local authorities might be given significant regulatory powers, while the central authority would retain those of a more limited scope or be in charge of setting the guiding principles and policies.

Rarely, governments may have explicit policies on Biosecurity which will naturally guide the legislative strategy to implement a Biosecurity approach. Other policies, although not expressly referring to Biosecurity, may have an impact on the legislative strategy. These would include the overall agricultural policy as well as policies regarding the environment, land use and trade. Good governance policies, such as access to information, participation

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13 J. Vapnek and D. Manzella, supra note 3, p. 10.
in decision-making, transparency and accountability of regulatory authorities will also affect the legislative design, as will policies on government structure and reform. For instance, if government policy does not encourage the establishment of new institutions, the legislative strategy will focus on establishing coordinating mechanisms to oversee existing institutions working in Biosecurity rather than proposing a new Biosecurity agency.

A Biosecurity approach puts traditionally sectoral institutions into a broader scheme of efficient management of risks to food and agriculture. It calls for inter-institutional cooperation and integration of functions, requiring strong commitment by the entire government and not only by individual ministries or agencies. Ideally, a Biosecurity approach will be implemented through comprehensive legislation, formulated in a participatory fashion, making it strong enough to change existing institutions and institutional behaviours.

VI. IMPLEMENT THE STRATEGY THROUGH NEW OR AMENDED LEGISLATION

The final step will be to identify the legislation that will be needed to implement the changes agreed upon by government policy- and law-makers, and then to prepare that legislation. Legislation will be needed toward two ends: first, to make substantive changes in the Biosecurity sectors, and second to change institutions or create new ones. For the first task, an example is where the plant protection law is outdated, an updated text will be needed to permit the government to carry out Biosecurity effectively with respect to phytosanitary issues. Or as another example, the absence of a biosafety law will call for the preparation of a new draft. If the assessment of sectoral legislation has revealed substantive deficiencies in several regulatory areas, amendments to the different pieces of legislation or the preparation of new legislation in those areas can be folded into a comprehensive legislative package and tabled before the legislative bodies.

Legislative change to empower new institutions or modify the functions of existing ones is the second area for action. As noted earlier, this may require the creation of a full-fledged Biosecurity agency or the establishment of a coordinating mechanism – although in some cases, governments may opt for some combination of the two, possibly to foster increased coordination as an intermediate step on the way to a fully autonomous Biosecurity agency. And although legislation may very well be needed, in some national contexts the development of memoranda of understanding between ministerial entities or
autonomous agencies may be sufficient and may be a quick and practical solution to institutional overlaps and gaps. Though without any legally binding force, these memoranda may establish a good working relationship between institutions and pave the way for coordinated action.

VII. CONCLUSION

The step-by-step methodology can be summarized as follows:

| A. | Analyse the legal framework covering the subject areas of Biosecurity |
|    | A.1 Collect and classify the legislation according to its level (parliamentary or subsidiary) and subject area (food safety, animal health, plant health, IAS, biosafety) |
|    | A.2 Evaluate the legislation |
|    | A.2.1 Find gaps in the substance of the sectoral laws |
|    | A.2.2 Determine compliance with international standards |
| B. | Review the mandates and functions of the various institutions responsible for Biosecurity controls |
|    | B.1 Identify the legal status of the institutions |
|    | B.2 Examine the legislated mandate of the institutions and compare it with the effective mandate |
|    | B.3 Identify gaps and overlaps among the activities of the institutions |
|    | B.4 Assess de facto arrangements to correct the gaps and overlaps of point B.3 |
| C. | Consider the feasibility of creating a new agency or establishing a coordination mechanism to implement a government-wide Biosecurity approach |
| D. | Elaborate a legislative strategy to pursue a Biosecurity approach |
|    | D.1 Consider the local context |
|    | D.2 Identify the legislative changes needed |
| E. | Implement the strategy through new or amended legislation |
|    | E.1 Elaborate legislation on substantive Biosecurity sectors |
|    | E.2 Prepare legislation to implement institutional change with regard to point C |
The methodology will naturally result in different recommendations for different countries. *Biosecurity* does not carry a one-size-fits-all solution; rather, it can be achieved in a variety of ways. The main objective is to provide countries with:

- a detailed understanding of the strengths and weaknesses of the legal and institutional framework for *Biosecurity*;
- plans for the implementation of necessary legislative and institutional change;
- an enabling framework of laws and regulations for the implementation of core Biosecurity functions in line with international legal requirements; and
- an organization or system with the mandate to perform controls and manage biological risks in food and agriculture.

The key is to choose solutions which are suitable for the time, the place, the policy context and the legal system of the country.
REFERENCES


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I. INTRODUCTION

This chapter contains an assessment of the existing system of Biosecurity in Ethiopia and its ability to accommodate international standards on plant and animal health, as well as food safety requirements. After this introduction, the chapter gives an overview of the legal and policy foundations that underpin the development of the Biosecurity laws and policies in the country.

In Ethiopia, there are some attempts to adopt specific regulatory provisions in scattered sectoral laws to realize the demands under each of the relevant international instruments. The National Biosafety Framework, which is currently being developed, is the only comprehensive document that focuses on one of the international agreements, the Cartagena Protocol. It has a comprehensive vision of biosafety and has provided the impetus for the drafting of a biosafety law and a number of directives.

This chapter also examines the mandates of regulatory institutions involved in the development and issuance of standards, such as the Drug Administration and Control Authority, the Ministry of Health, the Ministry of Agriculture and Rural Development, the Quality and Standards Authority of Ethiopia, the Ethiopian Health and Nutrition Research Institute, the Environmental Protection Authority and the Customs Authority.

Various laws existing in the country have empowered these agencies to undertake inspections of food quality, issue procedures and standards against risks of plant and animal diseases and thereby enhance human health and environmental sustainability. But as the laws were not initiated in a coordinated fashion and also owing to the low level of awareness on matters related to Biosecurity, the institutions are not functioning up to expectations. Some of the agencies, as will be seen below, lack either the necessary mandate or the requisite sectoral integration and coordination. This chapter analyses the exact lacunae in their mandates or the points of overlap in their functions.

The chapter goes on to outline the legislation in force and draft legal texts relevant to Biosecurity. The chapter discusses laws on seeds, plant protection and quarantine, pesticides registration, animal disease control and food

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1 This instrument is discussed in Chapter 2, Part VII.
safety, in particular in the areas where they are relevant to Biosecurity. There are gaps in the existing laws which exist either because of the complex law-making process in the country and lack of appropriate law-making capacity or the reluctance to adopt draft texts through the appropriate law-making channels. The chapter concludes by indicating the possible way forward in order to foster Biosecurity in Ethiopia.

II. POLICY AND LEGAL FOUNDATIONS OF BIOSECURITY IN ETHIOPIA

2.1. Constitutional provisions

The incorporation of environmental right provisions into the Constitution is recent in Ethiopia. In 1995, the Constitution introduced environmental rights as fundamental and inalienable to the people. There are no provisions in the Constitution that are directed at food safety or animal and plant health. However, some of the human rights provisions can be construed as incorporating the basic tenets of Biosecurity. The right to a "clean and healthy environment" is one of the rights that Ethiopian citizens are accorded as part of the fundamental and inalienable human rights (art. 44). What constitutes a clean and healthy environment is not explained in the Constitution. But a healthy environment requires protection of flora and fauna from organisms, chemicals, pests and invasive species. A clean and healthy environment cannot be ensured where minimum requirements of plant and animal health are absent. Thus, the protection of the environment against harmful substances or practices stems from the construction of these constitutional provisions.

A corresponding duty is imposed on the government to refrain from negatively affecting the health and development rights of the people (art. 92) and to promote those rights by issuing relevant protection schemes. All actors (state agents and non-state actors alike) shall respect the constitutional safeguards that are in place to ensure the balance between economic development and environmental protection (art. 43). The Constitution also provides for the improvement of the livelihood of the people of Ethiopia. Ethiopians also enjoy a right to be consulted on the adoption of policies and the implementation of

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projects affecting their communities. Prior informed consent of those communities is a pre-condition to the implementation of such projects.

Ethiopian citizens also have a right to be protected from undue displacement from areas where they live. In the event that this is compulsory (for instance, in case of health emergencies), people are entitled to monetary or non-monetary compensation, including relocation with adequate state assistance.

As stated above, these constitutional provisions are not specific to food safety or the protection of animal and plant health. However, they do lay down the basic conceptual framework for the setting of Biosecurity norms in the sectors of human health, environment and plant and animal health.

2.2. Policy coverage

2.2.1. Environment

In addition to incorporation of environmental issues in the Constitution, the framework of environmental protection in Ethiopia involves the formulation of an overarching environmental policy. The policy outlines principles to be followed in order to ensure the respect for environmental values, taking into account the economic, social and cultural circumstances of the country. The policy provisions relevant for Biosecurity in Ethiopia are discussed below.

The Environment Policy of Ethiopia (EPE) was approved by the Council of Ministers in 1997. The overall EPE goal is "to improve and enhance the health and quality of life of all Ethiopians and to promote social and economic development through the sound management and use of natural, human made and cultural resources and the environment as a whole so as to meet the needs of the present generation without compromising the ability of future generations to meet their own needs".3

In the EPE goal, there are features pertinent to the enhancement of human health and the protection of animals and plants from pests and diseases. For one thing, the policy targets as an ultimate goal the protection of the health and quality of life of the people. Though this goal does not provide for a list of the activities identified as harmful to

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human health, it can be inferred from the specific policy provisions that some elements of *Biosecurity* are instrumental to achieving the goal.

Principles of intra- and intergenerational equity are echoed in the policy in the sense that Ethiopian nationals have the right to utilize available natural resources, while at the same time they have the duty to conserve them for the use of future generations. Conservation of biodiversity and ecosystems appears in the policy. The policy also prohibits causing harmful and irreversible consequences to the natural and cultural heritage of the country.

The EPE contains sectoral and cross-sectoral elements that are of significance to *Biosecurity*. Under the sectoral policies, the most relevant aspects are those dealing with genetic, species and ecosystem biodiversity; human settlements, urban environments and environmental health; control of hazardous materials; and cultural and natural heritage. At the cross-sectoral level, EPE tries to link thematic issues like environment and population, community participation in decision-making, tenure and access rights to land and natural resources, and the importance of environmental impact assessment (EIA) and community participation in decision-making. EIA has a particular significance to ensure that the Ethiopian people and environment are safeguarded from alien elements that negatively affect the food system, ecosystems or any component of the environment. Owing to the importance of the EIA tool to a *Biosecurity* approach, this will be explored at more length later in this chapter.

The EPE envisages measures to develop and disseminate sustainable technologies to enhance agricultural production. This section of the policy can be the basis for regulating products of modern biotechnology under the draft National Biosafety Framework, particularly as regards the intentional release of such products into the environment.

There is a policy provision stating that ecosystems should be safeguarded from possible biological contamination through quarantine legislation. The possibility that some animals or plants may be infected with diseases and pests is also articulated in the policy for future action.

The EPE urges actions for the restriction of exotic species from biodiversity hotspot areas, thereby limiting the spread of some potentially invasive plants. Though the country does not have a stand-alone policy or specific legislation on invasive alien species, this policy element can be used as a basis for future
actions. The possible adverse effects of invasive alien species on biodiversity are also recognized under the water resources conservation section of the EPE. Its objective is to ensure that any proposed introduction of exotic species into water ecosystems is subject to detailed ecological studies and EIA. It also recognizes that natural ecosystems, particularly wetlands and upstream forests, are fundamental to rendering ecosystem services and hence deserve conservation. As with invasive alien species, despite this policy statement there is no law in place governing conservation and utilization of fisheries resources.

The policy goals laid down in the EPE seem to reflect the government's commitment to conserve natural resources and protect the environment. However, it is clear that this commitment has to be substantiated through detailed and enforceable rules. The EPE has a mechanism for its periodic revision, although no initiative has been taken in that respect after the adoption of the policy.

In spite of the policy foundation, the quarantine laws of the country are far from meeting international standards. The problems emanating from the movement into and out of the country of organisms that can be categorized as plant pests and animal diseases remain without an adequate legislative response.

2.2.2. Biodiversity

Ethiopia has a national policy on biodiversity and research which was adopted in April 1998. The objectives of this policy are to ensure that genetic resources and ecosystems as a whole are conserved, developed, managed and sustainably utilized. Biodiversity conservation and development programmes should be duly integrated into the country's agricultural, health, industrial and overall national economic development strategies and plans. Promoting regional and international cooperation in biodiversity conservation, development and sustainable use is also another policy goal.

Some elements of the biodiversity policy can be construed to encompass the environmental aspects of Biosecurity (in particular, the loss of genetic diversity that may result from pests and diseases). The policy was initiated under the auspices of the Institute of Biodiversity Conservation and Research (IBCR), now the Institute of Biodiversity Conservation. The mandate to carry out research was removed and is now being undertaken by the Ethiopian Agricultural Research Institute, which seems to have assumed all tasks of research and development that used to be carried out by the IBCR. This
overlap in mandates has caused the research component of the policy to lose efficacy. These policies will be discussed later in this chapter with respect to their relevance to Biosecurity.

III. INSTITUTIONAL AND LEGAL REGIME OF BIOSECURITY IN ETHIOPIA

3.1. Institutional framework

3.1.1. Crop Protection Department

Ethiopia joined the International Plant Protection Convention (IPPC) on 20 June 1977. The Ministry of Agriculture is the responsible body vested with the power to coordinate all efforts as regards compliance with this instrument. This ministry was reorganized in 2004 and renamed the Ministry of Agriculture and Rural Development (MOARD). Its mandate includes all measures necessary to:

- conduct quarantine controls on plants, seeds, animals and animal products; and
- prevent outbreaks of animal diseases and plant pests.

The Crop Protection Department is the national plant protection organization (NPPO) of Ethiopia. It has the following three divisions:

a. *Pesticide Registration and Control Team.* The team is composed of a few experts mainly tasked with registering agricultural inputs. They operate under the Pesticide Registration Decree No. 20/1990 which regulates pesticide import permits;

b. *Crop Protection Laboratory and Quarantine Team.* This division is responsible for ensuring that all imported and exported agricultural products are inspected and verified as free of any injurious insects, pests, diseases and noxious weeds. With a federal mandate, the division oversees the functioning of a number of quarantine stations. The stations currently functioning are those at Bole Airport, Dire Dawa, Metema, Moyale and Nazareth (Central Rift

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4 The IPPC is discussed in Chapter 2 Part III.
Among these, it is only the Bole Airport and Moyale stations that carry out import inspections. The rest are performing only inspections for export;

c. **Crop Protection Division**: This division controls migratory pests such as African Army worm, locust, etc. It has a federal mandate as it looks only at transregional issues.

Inspectors are assigned to each of the quarantine stations listed above in paragraph b. Inspectors are empowered to search, inspect, analyse, treat and seize any infested or infected plants, plant products and articles. The inspectors are not properly trained and, being few, do not have the capacity to monitor all the movements of plants and plant materials across the borders. In most instances, inspection is primarily carried out through visual observation.

None of the quarantine facilities in Ethiopia is adequately equipped. With the exception of those at Moyale and Nazareth, none has any laboratory facilities. Most of the stations also lack basic facilities such as greenhouses or fumigation centres.

Some laboratory equipment is found in the different agricultural research institutions. For instance, the Holeta and Melkassa Agricultural Research Centres have independent laboratory facilities. The Ethiopian Agricultural Research Institute has a laboratory facility that undertakes molecular marker techniques to characterize local poultry. The Sebetta Laboratory is serving as the National Animal Health Diagnostic Centre and has laboratory equipment for this purpose. Some of the science universities in the country such as the Biology Department of Addis Ababa University and Haromaya University have laboratories destined to fulfil the research needs of their students. To cope with these capacity constraints, it has been proposed to establish a central laboratory to employ equipment scattered in the many sectoral agencies.

The lack of capacity is also accompanied by institutional conflicts. With regard to plant health, there is a conflict between the Crop Protection Department of MOARD and the IBC. The latter is the national institution established in 1976 with responsibility for coordinating efforts to stop the rate of genetic erosion by promoting conservation activities. It is vested by
law with the responsibility of granting access to genetic resources under certain conditions. Although the ambit of its authorizing law is access to genetic resources and not plant health, all plant material ready for export is interpreted as access granted on the germplasm, thus requiring a permit from the IBC. This often leads to a conflict between the operating procedures of the two entities. A memorandum of understanding or a legislative instrument would go far to resolving the conflict.

There is no official quarantine pest list kept within the Crop Protection Department. Nor has there been so far any attempt to designate a pest free area in Ethiopia. Regarding pest risk analysis, there are no defined or elaborate procedures that would enable the country to comply with the demands of the IPPC. Most of the international standards developed under the IPPC are not being implemented in the country.

3.1.2. Animal and Fisheries Resources Development and Regulatory Department

The Animal and Fisheries Resources Development and Regulatory Department within MOARD is currently empowered to undertake regulatory functions relating to livestock development in the country. Animal and animal products are inspected by this department. There are four animal quarantine stations in the country, namely, at Afar, Dire Dawa, Nazareth and Bole Airport. There are veterinarians, senior inspectors and junior inspectors assigned at each of these posts. At the veterinary inspection post of Bole Airport (which is the main port of exit for animal products via air shipment), two veterinarians and four senior inspectors are assigned to inspection in order to control the movement of animal products. Apart from these there is a national animal health diagnostic centre (Sebetta Laboratory) in the town of Sebetta which is in the vicinity of the capital city.

There is generally a lack of integration in terms of plant and animal inspection activities at the ports of entry and exit. Apparently, the only quarantine station that is common both to plants and animals is the post at

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5 Proclamation on Access to Genetic Resources and Community Knowledge No. 82/2006. This proclamation prescribes procedures for facilitating access by foreign users to the plant and animal resources of Ethiopia. It also provides an institutional mechanism for equitable sharing of the benefits arising from their utilization.
the Bole Airport. Laboratory and inspection facilities are also scattered, which leads to mismanagement.

Within the Animal and Fisheries Resources Development and Regulatory Department, the Veterinary Services Team is responsible for maintaining animal health and the safety of food products of animal origin. There is, however, a serious constraint to the efficient functioning of the team. When compared to the overall importance of the livestock sector, particularly to the livelihoods of the nomadic community living in the Afar and Somali regions, the staffing levels within the department are too low\(^6\) and personnel often lack the requisite skills. It is often commented that the country has the lowest animal health care coverage in the entire sub-Saharan community\(^7\).

The OIE Terrestrial Animal Health Code specifies the guidelines for safe animal and animal product trade.\(^8\) These guidelines specify that livestock products must originate from countries or specified geographical areas (zones) of a country that are free from major animal diseases capable of causing economic losses or human diseases. In compliance with these requirements, Ethiopia attempted to designate a disease free zone covering the regions of Afar, Borena and Ogaden with a view, among other things, to maximizing profits from the rich livestock resources in these parts of the country.\(^9\) The designation has not proven to be effective as the community is mainly made up of nomadic pastoralists and diseases will not be contained outside of the disease free zone as intended. Currently, there is no disease free zone officially communicated to the OIE.

\(^{6}\) See *Ethiopia Trade and Transformation: Synthesis, Diagnostic Trade Integration Study*, available at [www.integratedframework.org](http://www.integratedframework.org), according to which this veterinary services team consists of only eight professionals and a total of about 500 veterinarians, 800 para-veterinarians and 3,400 animal technicians. These experts manage a network of about 930 clinics, 650 animal health posts and ten diagnostic laboratories. The private veterinary sector is weak, with a total of 57 private veterinarians, 64 private clinics and about 150 private animal health assistants.

\(^{7}\) See id.

\(^{8}\) The OIE, or World Organization for Animal Health, is discussed in Chapter 2, Part IV.

3.1.3. Environmental Protection Authority

The Environmental Protection Authority (EPA) is an autonomous public institution of the federal government. It has the role of coordinator among environmental protection agencies at the federal and regional levels. EPA formulates policies, strategies, laws and standards for the environment and enforces them. Article 6 of the proclamation establishing the EPA\textsuperscript{10} includes the powers and duties to:

- coordinate measures to ensure that the environmental objectives provided under the Constitution and the basic principles set out in the environmental policy of Ethiopia are realized;
- prepare environmental policies, strategies and laws and upon approval, monitor and enforce their implementation;
- establish a system of environmental impact assessment (EIA);
- review EIA reports of such projects and notify its decision to the concerned licensing agency, and audit and regulate their implementation;
- set environmental standards and ensure compliance with them;
- formulate policies, strategies, laws and programmes to implement international environmental agreements to which Ethiopia is a party; and
- coordinate, promote and as may be appropriate, carry out research on environmental protection.

EPA operates at the federal level while all eleven regions of the country (including the autonomous cities of Addis Ababa and Dire Dawa) have established respective environmental bureaux to enforce environmental standards.\textsuperscript{11} Though all the core responsibilities listed under the provisions of this law are not specific to Biosecurity, they have a bearing on aspects of it. The term "environment", as defined in Proclamation No. 295, in itself

\textsuperscript{10} EPA Establishment Proclamation No. 9/1995. A proclamation is an act of Parliament while a regulation is a pronouncement of the Council of Ministers. A directive on the other hand is a subsidiary instrument that is adopted by the competent executive agency.

\textsuperscript{11} Proclamation No. 295/2002 (Proclamation to provide for the Establishment of Environmental Protection Organs). Under article 15(2), regional states are responsible for ensuring the implementation of federal environmental standards or, as may be appropriate, issuing and implementing their own no less stringent standards.
embodies elements of protection of human, animal, and plant life. According to article 2(3), it consists of:

"the totality of all materials whether in their natural state or modified or changed by humans, their external spaces and the interactions which affect their quality or quantity and the welfare of human or other living beings, including but not restricted to land, atmosphere, weather and climate, water, living things, sound, odour, taste, social factors, and aesthetics."

The EPA has initiated the drafting of a number of laws. The EIA system operational in Ethiopia is one of the creations of this institution and is a key tool in achieving environmental sustainability and the enhancement of animal and plant health. The law on EIA and its implementation practice will be discussed later in this chapter.\textsuperscript{12}

3.1.4. Drug Administration and Control Authority

The Drug Administration and Control Authority (DACA) was established as a semi-autonomous regulatory agency through Proclamation No. 176/1999 with the objective of ensuring safety, efficiency and quality of drugs and regulating their production, distribution and use. DACA has the power to set standards and ensure their observance, to control the quality of raw materials and packaging and to monitor drugs and set standards for traditional medicine practitioners and users. Along with these functions, inspectors are empowered to search any premises, conduct inspections, seize documents and take samples of materials. DACA performs import and distribution controls through a system of registration and import permits.

3.1.5. Ethiopian Health and Nutrition Research Institute

The Ethiopian Health and Nutrition Research Institute (EHNRI) was established by Council of Ministers Regulations No. 4/1996. The major objectives of the institute are:

\textsuperscript{12} See Section 3.2.9.
• to conduct research on the causes and spread of diseases, on nutrition, on traditional medicines and medical practices and on modern drugs;
• to support activities for the improvement of health in the country; and
• to contribute to the development of health science and technology.

Apart from general research duties, EHNRI is vested with powers and duties that are relevant to food safety. It is, for instance, expected to undertake studies on the causes, health impacts and distribution of food-borne diseases and conduct nutritional science and technology research on food items.

3.1.6. Ministry of Health

The Ministry of Health (MOH) is one of the executive organs of the government established through Proclamation No. 4/1995. The mandate of MOH, as outlined under article 22 of the proclamation, includes:

• determining standards to be maintained by health services;
• determining the required qualifications of professionals engaging in public health services at various levels;
• devising and following up the implementation of ways and means of preventing and eradicating communicable diseases;
• undertaking necessary quarantine controls to protect public health; and
• undertaking studies with a view to determining the nutritional value of foods.

The responsibilities of MOH regarding food safety and protection of public health are further elaborated in the Public Health Proclamation (Proclamation No. 200/2002). The goals of this proclamation as stated in its preamble are:

• to bring about attitudinal change in society through the primary health care approach with a view to solving most of the health problems of the country; and
• to promote the health of the society for future generations.
In order to achieve these objectives, the proclamation has provided for specific rules on food quality control and provisions on applicable food standards in the country. According to article 2(1), food quality shall be ensured through compliance with biological, chemical and physical standards set nationally and internationally.

A number of important terms are defined in the same article. "Food" is defined to mean any substance whether processed, semi-processed or raw which is intended for human consumption and includes drinks, chewing gum and any substance which has been used in the manufacture, preparation or treatment of food, but it does not include tobacco, cosmetics or substances used only as drugs. 13 "Food additive" is a substance added to food to improve its taste, colour, preservation or appearance and which is considered to become a component of food. Minimum requirements of food quality are set under articles 8 and 9 of this proclamation which are discussed at length later in this chapter.

In order to ensure compliance with the law, inspectors are assigned duties and responsibilities such as the power to:

i. enter and inspect any premise where the inspector has sufficient reason to believe that there exists a situation endangering public health;

ii. seize any article or material which is the result of any act committed contrary to law or used for the commission of the illegal act or has any connection with the commission of the illegal act;

iii. order that the premises remain closed for a limited period of time;

iv. take, where necessary, samples of articles, materials or goods from any premises or building, or any sample of air from within the premise or from the compound;

v. cause the destruction of articles, materials or goods found in any premises or building where there is sufficient reason to believe that such goods are dangerous to health, or that they cause or can cause another danger;

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13 This mainly tracks the Codex definition.
vi. request any information from any person which the inspector believes can give any information relevant for the investigation; and

vii. cause the institution of prosecution by the authorized organ.

3.1.7. Quality and Standards Authority of Ethiopia

The Quality and Standards Authority of Ethiopia (QSAE) was first established in 1970 as the National Standard Body. In 1998, it was re-established through Proclamation No. 102/1998, which was later amended by Proclamation No. 413/2004.

The QSAE is an autonomous federal organ operating under the Ministry of Trade and Industry (MOTI), whose major task is to approve Ethiopian product standards, including for crops and animal products. The modalities of operation of the institution and the main elements of its standard-setting activity are discussed later in this chapter.14

3.1.8. Institute of Biodiversity Conservation

As noted earlier, the Institute of Biodiversity Conservation and Research was established in 1976 and underwent gradual evolution, with its mandate and name undergoing frequent change. Initially, it was established as the Plant Genetic Resources Centre of Ethiopia and was focused on the collection and ex situ conservation of crop plants with high research and economic importance. The centre gave priority to crop types facing immediate danger of extinction and genetic erosion.

The institute was renamed the Biological Diversity Institute in 1994 and was tasked with, among other functions, conserving genetic resources and providing germplasm for the improvement of crops, acquiring new germplasm from other countries, documenting indigenous community knowledge and establishing field gene banks and botanical gardens.

In 1998, it was renamed the Institute of Biodiversity Conservation and Research (IBCR) through Proclamation No. 120/1998. Under this proclamation the institute has the power to initiate policy and legislative proposals and survey genetic diversity and distribution. It can also undertake

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14 See Section 3.2.8.
ex situ and in situ activities to conserve biological resources and implement international conventions, agreements and obligations on biodiversity. It is now the focal point for the Convention on Biological Diversity (CBD).15

The IBCR was restructured by Proclamation No. 381/2004 into a semi-autonomous entity under MOARD and assumed its current name – the Institute of Biodiversity Conservation (IBC). The IBC has various powers and duties relevant to Biosecurity and in particular, its environmental aspects. These include:

- cooperating with the concerned federal and regional authorities with respect to protection of biodiversity;
- encouraging and supporting public participation in the conservation, development and use of biological resources; and
- developing regional and international cooperation on biodiversity conservation and research activities, based on international agreements and national legislation.

3.2. Legal framework

3.2.1. Plant protection

MOARD is given the mandate for plant health by virtue of Council of Ministers Regulation No. 4/1995. Under this law MOARD shall assume the duty to establish quarantine of plants to prevent the spread of plant pests and to regulate the movement of plants, plant products or other articles into or from a specified area. Once the plant species are identified and the infested or infected materials isolated, MOARD can also treat it or in the worst case scenario dispose of it at the expense of the owner. This process limits the possible damage that would result from inappropriate disposal of the infected plants or plant products.

MOARD is further empowered to restrict the importation of certain plants which do not have import permits and phytosanitary certificates duly issued by the plant protection authorities of the exporting countries. MOARD has the responsibility for issuing phytosanitary certificates for export of Ethiopian plants and plant products.

15 This convention is discussed in Chapter 2, Part VI.
The Plant Quarantine Regulations (Council of Ministers Regulations No. 4/1992) further detail the provisions on import and export. The regulations prescribe that any plant or other articles, premises or conveyances found infected shall be treated or destroyed, as the case may be. Quarantine controls and documentary verification of phytosanitary certificates on all imported plants are required. Some plant species are prohibited from entering the country. For others there are restrictions and a permit should be secured from MOARD before they may be imported. The regulations also provide for the declaration of quarantine areas and subsequent control measures.

The regulations are silent with regard to surveillance of pests within the country. They contain a list of quarantine pests which is not updated and includes pests that have been endemic in the country for a long time. For instance, parthenium is a weed which is endemic to South and North America but is reported to have been noticed in Dire Dawa, Hararghe and eastern Ethiopia as from 1988. It was supposedly imported from subtropical North America "as a contaminant of grain food aid during the 1980s famine" and subsequently spread in the country.

3.2.2. Seed and variety release

According to its preamble, the Seed Proclamation is intended to achieve increased crop production by enabling farmers to use high quality seeds, particularly for improved varieties, and making those seeds available on the market. The proclamation does not directly deal with plant health. However, article 14 provides that any seed produced and processed locally, imported or exported shall be of a variety registered by the National Agricultural Inputs Authority (NAIA) and shall conform to the requirements and standards of the country.

Phytosanitary inspections of seeds are undertaken before the variety is released. However, as seed is also one of the plant parts subject to the quarantine regulations, there is often a conflict in mandate between the

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17 Id.
National Agricultural Inputs Department of MOARD, which replaced the NAIA as Ethiopia's seed authority, and the Crop Protection Department.

There is a de facto legal instrument\textsuperscript{19} that puts in place the applicable procedure regarding seed development and release in the country. Under this procedure, the National Variety Release Committee approves the release of hybrids and varieties developed by governmental and private institutions, makes the necessary arrangements to conduct quality tests in collaboration with other agencies and registers the released varieties and hybrids.

3.2.3. Agricultural inputs

NAIA was established under Proclamation No. 288/2002 with the mandate of regulating agricultural inputs (plant seeds, fertilizers and pesticides). NAIA was established with the basic purpose of ensuring increased production and productivity of the agricultural sector. The restructuring of the Ministry of Agriculture has eliminated the autonomy of the NAIA and made it a department within MOARD. The Agricultural Inputs Department of MOARD has the task of ensuring the health of all agricultural inputs. The department is mandated to inspect the quality of seeds, fertilizers and pesticides to be released on the local market or those to be exported or imported.

3.2.4. Pesticides

The pesticide regulatory instrument is a Council of Ministers decree enacted in 1990 (Regulation on Pesticide Registration, Council of Ministers Special Decree No. 20/1990). It lays down a scheme of control to minimize the adverse effects of pesticides on human beings, animals, plants and the environment. "Pesticide" is defined as any substance or mixture of substances intended to prevent, destroy or control any pest, including vectors of human or animal disease, unwanted species of plants or animals causing harm during or otherwise interfering with the production, processing, storage, transport or marketing of food, agricultural commodities, wood and wood products, animal foodstuffs or substances

\textsuperscript{19} National Variety Release Procedures and Mechanisms, adopted on November 2001 by the National Authority for Inputs in Agriculture. These procedures and mechanisms are in place and have been used by the National Variety Release Committee for over a decade without any challenge to their authority by individuals or legal entities.
which may be administered to animals for the control or insects, arachnids or other pests in or on their bodies. The decree prohibits the manufacture, import, sale and use of unregistered pesticides in Ethiopia.

There are no official standards or criteria nationally adopted to regulate the registration process in cases of import. There also appears to be a mandate conflict on the registration of veterinary drugs for import. These drugs are perceived by MOARD to be included within the wider definition of pesticides, whereas DACA considers them no different from human drugs and claims competence to regulate them. An inter-ministerial committee was assigned the responsibility to resolve the conflict. Currently, the registration of all forms of veterinary drugs is performed by DACA, which issues import licences. It also keeps a list of registered veterinary drugs and registration guidelines. There is a perceived need to upgrade the status of this list to be an enforceable law.

3.2.5. Animal health

Animal disease control is dealt with under Proclamation No. 117/1998. The Animal Marketing Authority (AMA) was established in the proclamation with the objective of promoting the domestic and export marketing of animal products and by-products through increasing support and improving quality. In order to achieve these goals, the AMA has the power to issue quality control directives on exportable or importable animals, animal products and by-products and to follow-up on trading activity. It also has to ensure that the exportable items meet international standards.

The AMA has been dissolved and there is no later authority assigned to take over its function. The activities it used to undertake are now being followed up by the Animal and Fisheries Resources Development and Regulatory Department within MOARD.

In 2003, the Ethiopian Government designed an export development strategy which gives particular attention to the promotion of labour-intensive production and processing for export. Meat, livestock, hides and skins are priority export commodities within this strategy. In order to win on the competitive international market, the government is building capacity to comply with the different international standards, particularly those emanating from the OIE.
Legal instruments regulating animal diseases, livestock development and meat inspection are summarized hereunder.

The *Animal Disease Control Proclamation No. 267/2002* mainly deals with the prevention and control of diseases; outbreak notification; establishment of quarantine stations; entrance and exit ports for export of livestock and livestock products; international animal health sanitary certification; and animal movement permits. MOARD is authorized by this proclamation to declare by public media an area infected by a notifiable animal disease (art. 4). This notice should specifically indicate the infected area, the type of disease and the measures to be taken. Specific actions which can be taken in relation to an infected area are set out in article 5. MOARD is also assigned the mandate to extend, diminish or otherwise alter the limits of a declared infected area.

The law also incorporates provisions stating the conditions under which animals, their products or by-products are exported. Article 12 states the requirement to keep animals for export in a quarantine station for a specified period. Animals for export shall also originate from an area that is free from notifiable animal diseases and be accompanied by a movement permit. Persons who transport export animals, animal products and by-products are required to comply with transport requirements and use designated exit posts.

The importation of animals and their products should follow similar requirements under article 13 of the law. The importer should obtain an entrance permit by applying to MOARD stating the type of product, country of origin, quantity, means of transport, date of arrival, port of entry and transit countries prior to importation of animals, animal products and by-products. Imported items should be checked at the port of entry by an animal health officer.

There is also a provision on animal movement and the requirement of a permit for such purposes. Article 14 states the duty to obtain an animal movement permit from the animal’s place of origin in order to transport animals from woreda to woreda or from region to region. The animal movement permit should indicate the animal's place of origin, destination, route, type and number of animals, health status and other necessary details.

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20 A "woreda" is a lower administrative unit while a "kebele" is the lowest.
The Meat Inspection Proclamation No. 274/1970 confers a mandate on MOARD to control and regulate the production, processing and handling of livestock products.

The Meat Inspection Amendment Proclamation No. 81/1976 mandates MOARD to issue regulations and establish criteria for livestock production for human consumption, including classification of products and inspection of processing facilities.

The Meat Inspection Regulations No. 428/197 are issued under Proclamation No. 274. They establish the requirements for setting up abattoirs and commercial establishments dealing with slaughter, preparation and processing of livestock products for export from or import into Ethiopia.

There are also a series of draft legislative instruments:

The Draft Regulations for Animal Diseases Prevention and Control set rules for disease reporting, investigation and surveillance mechanisms at federal and regional levels. They also set the modalities for the control of disease outbreaks.

The Draft Regulations for Controlling Movement of Animals and Transportation of Animal Products and By-products set the mechanisms to prevent the spread of infectious diseases.

In addition to this, a series of guidelines, which are non-binding instruments issued by MOARD, have been prepared for:

Meat inspection, hygiene and construction of export abattoirs: These guidelines were approved in 2000 to set the standards of good practice.

Operational procedures of export abattoirs: These guidelines set out the procedures for examining animals destined for slaughter and for implementing sanitary measures in abattoirs.

3.2.6. Food safety

In Ethiopia, the leading government institutions responsible for food safety include MOH, MOARD, the QSAE, MOTI and the Ethiopian Manufacturing Industries Association. These institutions work together in
organizing training workshops, setting standards and drafting regulations. Since 2002, these bodies have established a Technical Committee that implements food safety assurance systems in accordance with the international market requirements, supported by the United Nations Industrial Development Organization.

Although there is no comprehensive food safety policy in the country, safeguarding the public from communicable and infectious diseases is clearly addressed in different policies such as the National Health Policy that gives due emphasis to prevention and control of the major health problems of the country, which would include problems arising from food safety.

As noted in Section 3.1.6., the Government of Ethiopia has issued Public Health Proclamation No. 200/2002 which gives general powers on public health matters to MOH. The proclamation establishes an advisory board at the federal level and regional health bureaux at the regional level for the purpose of advising MOH on the implementation of the proclamation. This task inherently includes food safety since it is one of the important factors in ensuring public health.

There are minimum requirements set under this law to ensure national food quality control (art. 8). Accordingly:

- it is prohibited to prepare, import, distribute or make available to consumers any food which is unhygienic, contaminated, unwholesome or mislabelled and does not meet the standards of food quality;
- any food intended for human consumption should meet the standards of food quality and be labelled and preserved in a healthy manner;
- any person who produces or distributes salt for human consumption shall ensure that it meets the standard requirement of iodine content; and
- no person shall use any testing laboratory unless it is registered by the health authorities.

It is a requirement under the law that a person engaged in any activity of selling, producing for sale, storing, preparing or preserving food intended for human consumption should meet the standards set by MOH. Regarding
water quality control, it is prohibited to import, produce or distribute to the public bottled mineral water or plain water unless its quality is verified.

Based on this proclamation, draft food safety regulations are completed and awaiting endorsement by the Council of Ministers. Most regional health bureaux have enacted their own regulations that fit their regional context based on this proclamation.

Committees consisting of government representatives, food manufacturers, food traders, food scientists, food inspectors, medical and veterinary experts, consumers and other stakeholders set food standards. These standards can be company, national, regional or international depending on the scope of their application. The QSAE has developed about 450 food-related standards, most of which have been implemented and have been made technical regulations by Regulation No. 13/1990. Standards are the technical basis for food safety inspection activities. Food products are inspected and controlled through third-party certification schemes.

A number of institutions are assigned, through the proclamations establishing them, to undertake food safety inspections in the country, including MOH, QSAE, EHNRI, DACA and the Customs Authority. There are overlaps in the exercise of those functions that are hampering the effectiveness of controls. For instance under MOH Proclamation No. 4/1995, one of the functions of MOH as regards ensuring public health is undertaking quarantine controls. This, however, is a duty assigned to MOARD under other laws. The task of setting food quality standards, assigned to MOH, is also another problematic issue as it also falls within the mandate of QSAE. The latter is in fact in the process of developing food quality standards to be applied nationally.

Coordination of activities at the lower level of the hierarchy remains to be established and strengthened. Responsibilities and mandates are not clearly defined, demarcated or streamlined, resulting in insufficient coordination of activities, duplication of efforts, misuse of human resources and waste of meagre financial resources allocated to the sectors. In order to overcome these problems, the existing Technical Committee has established the National Food Safety Council whose members are drawn from regulatory bodies, research institutes, industry, consumers and higher learning institutes involved in food safety.
3.2.7. Biosafety

Ethiopia is a party to both the CBD and its Cartagena Protocol. In an attempt to implement the obligations under the protocol, the country implemented a UNEP/GEF-funded biosafety project which resulted in the National Biosafety Framework (NBF). The NBF includes a draft Biosafety Proclamation.

According to the preamble, the objective of this draft proclamation is to enable the country to benefit from the advantages of modern biotechnology, by managing the possible risks occasioned as a result of the application of the technology on human and animal health, biological diversity and the environment. The precautionary principle, which requires the employment of cost-effective safety actions to prevent potential harm even in the absence of conclusive scientific evidence, underpins the draft proclamation.

The draft proclamation establishes procedures of prior notification to and authorization by the EPA for research and development, import, export, transit, handling, contained use, transport, placing on the market, use as a pharmaceutical for humans or animals, use as food, feed or for processing of any genetically modified organism (GMO) or products thereof. EPA is designated to be the responsible authority for the approval (or rejection) of applications. The proclamation initially envisaged a committee of experts from various regulatory agencies to advise the EPA but this concept has subsequently been abandoned.

The applicant is required to undertake risk assessment to identify potential risks of GMOs or derived products on human and animal health and biological diversity, including socio-economic conditions and cultural norms and the environment in general (art. 9). A GMO exporter is required to provide evidence of the advance informed agreement of the importing country. EPA is not precluded from taking more restrictive actions or prohibiting the intended export (art. 13). EPA is required to make any application available to the public and technical experts and solicit their comments.

The draft proclamation also provides for the identification, labelling and packaging of GMOs or their products, with the EPA tasked with establishing standards in this regard. Post-authorization monitoring and inspections are also regulated in the draft proclamation. Criminal sanctions
are imposed on offenders who contravene the mandatory obligations of the proclamation such as those on notification, risk assessment and compliance with standards.

The draft proclamation contains the basic provisions that are required in order to implement the obligations of the Cartagena Protocol. If approved, it would be an important tool within the national Biosecurity system to manage risks to the environment and human health arising from GMOs.

3.2.8. Product quality standards

As seen above, the QSAE currently sets standards for agricultural products and is empowered to ensure compliance with them. The standards are harmonized with the pertinent standards of other countries so that they will not constitute trade barriers under the World Trade Organization Agreement on Technical Barriers to Trade.21

To monitor compliance, the QSAE acts at three levels: local manufacturing, import and export. Any organization engaged in the food manufacturing business will give samples of the products to the laboratory of QSAE for inspection. Routine inspections on the premises are also conducted on a quarterly basis. In the case of imports, the QSAE issues pre-import authorizations and inspects at ports of entry, with sampling and laboratory testing if necessary.

For exports, the QSAE operates a permit system for food exporters. Inspections on food items for export are carried out mainly on oil seeds and pulses. The QSAE does not inspect the content of manufactured products but only verifies compliance with labelling requirements. The QSAE is also responsible for setting plant and animal health standards but their implementation is left to the responsible units within MOARD.

3.2.9. Environmental impact assessment

The Environmental Impact Assessment (EIA) Proclamation No. 299/2002 entered into force in December 2002. The proclamation tasks EPA with the elaboration of a series of projects and activities for which an EIA is required.

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21 This agreement is discussed in Chapter 2, Section 2.2.
It establishes criminal penalties in cases of false representations in an EIA report (art. 8). Though not specifically addressing Biosecurity issues, this law is used to manage environmental risks for imports of invasive plant and animal species. A person desiring to engage in such an import business is required to produce a clearance from the regional environmental agencies. It is only when an authorization is secured that an import is allowed (art. 3(3)).

The steps generally followed are that the applicant prepares an EIA study which is subsequently reviewed by the competent regional environmental agency. After review of the EIA report, the environmental agency will decide either to approve or refuse the proposed request. This decision is expected to be followed by the issuance of a licence or permit by the rendering agencies (art. 3(3)). These agencies should check the environmental clearance in advance of granting any form of permit or operational authorization.

The Crop Protection Department and the Animal Inspection Department of MOARD should follow this mandatory provision of the law. This, however, does not seem to be taking place. The ports of entry and exit that perform quarantine functions only examine the physical items subject to movement and do not require any environmental clearance for shipment to or from the country. This is an example of a lack of interagency cooperation. MOARD has its own recently established in-house Environmental Management Unit. This unit is mandated, under article 14 of Proclamation No. 295/2002, to ascertain that environmental standards are complied with.

The implementation of the EIA proclamation is further hampered by a number of other constraints. The list of projects and activities that need to undergo compulsory EIA procedures is not in place yet and, as a consequence, it is unclear whether a certain import requires EIA or not. This is one grey area that needs to be clarified. The procedures for undertaking EIA studies (including the qualifications of the expert that can undertake an EIA study or the composition of panels of experts) are not set forth in law. The existing law also provides that the EIA review process include participation of the affected communities (art. 14) but so far there is no record of such consultation undertaken. Moreover, the EPA reviewers of EIA studies often lack proper training.
IV. BIOSECURITY ASSESSMENT AND RECOMMENDATIONS

Although Biosecurity does not appear in any policy document in Ethiopia, the different sectoral policies of Ethiopia provide the necessary policy foundations for a Biosecurity approach. For instance, the EPE, although not in a detailed manner, provides for the need to control the introduction of pests or invasive alien species (IAS) into ecosystems. The existing policy provisions can be employed as the basis for the development of legislation. However, as regards other Biosecurity components, there is no on-going initiative for the upgrading of legislation. Sectoral legislation in the area of animal disease, plant pest control and quarantine regulations is not comprehensive. The laws generally are not harmonized with respect to the mandates of the implementing institutions. This often leads to conflict rather than cooperation.

Seen in this light, the overall legal system of Biosecurity in Ethiopia is only at a rudimentary level. It is full of gaps that render it incomplete compared to dynamic developments in the international arena. Many of the Biosecurity-related international instruments are not yet fully domesticated. For instance, as seen above there is no IAS law to give effect to the obligation of Ethiopia under article 8 of the CBD.

Draft legislation on biosafety has been prepared. However, the drafting of the biosafety proclamation as well as the National Biosafety Framework was the result of technical assistance under UNEP/GEF. After the assistance ended, the draft has not proceeded through the legislative process. A similar situation inheres with regard to other draft legislation of relevance to Biosecurity:

- draft Regulations for Animal Diseases Prevention and Control; and
- draft Regulations for Controlling the Movement of Animals and the Transportation of Animal Products and By–products.

These draft documents should be finalized and undergo the necessary approval process. One opportunity that these drafts avail is the possibility of incorporating Biosecurity concepts into them, such as the streamlining of import and export procedures or the pooling of resources to conduct risk assessment. In this regard, an entry point is the NBF.
The status of some legal instruments should also be revisited. Some Biosecurity instruments are mere guidelines that are subject only to voluntary compliance. These should be given more legal weight and be transformed into directives, which are enforceable but easier to approve than regulations or proclamations. Included in this category are the guidelines on meat inspection, hygiene and construction of export abattoirs.

The gaps in the EIA system in Ethiopia should be given attention as EIA is essential to control the damage that may be caused to flora and fauna as a result of imports. At present, the implementation of this important tool demands adopting subsidiary regulations to put the prescriptions of the law into action. For instance, as stated earlier, the activities and projects that require a full-scale EIA should be listed and a mechanism be devised to coordinate the working procedures of the environmental agencies especially at the ports of entry.

Most of the institutions engaged in the management of animal and plant health as well as food safety are constrained by conflicts and overlaps in mandate that seriously lessen their effectiveness. Biosecurity is not currently being handled by a single institution. Understandably, this is due to the multidisciplinary nature of the subject. However, the present situation has created lacunae in the implementation not only of international standards but also of the laws in force within the country. The handling and implementation of Biosecurity obligations is generally a task scattered within a wide array of agencies. In order to implement the food quality standards, plant life and animal health standards of the Codex Alimentarius, OIE and the IPPC, there is a need to improve this institutional infrastructure.

Owing to the sectoral orientation of the agencies handling Biosecurity in Ethiopia, interagency cooperation is very poor. This can be solved by the assignment of an oversight body at a higher level vested with the task of supervising the implementation of Biosecurity-related activities. An interagency committee composed of the major institutions involved (EPA, MOARD, MOH, EHNRI, DACA, QSAE) can be formed to address Biosecurity issues in the country. It is important to make such a committee accountable to a higher-level political authority such as the Prime Minister's office, to give it more power and efficacy.

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22 The Codex Alimentarius is discussed in Chapter 2, Part V.
The disease surveillance and quarantine system of the country is constrained because of the unwise use of limited resources such as laboratories, laboratory equipment and the necessary personnel. One urgent response would be to pool the country's meagre human and technological resources and establish a centralized system for risk assessment and risk management. This would also ensure the necessary synergies among different experts and the reliability of the risk analysis process. Legislation could be adopted in this regard to consolidate all the scattered surveillance and quarantine activities undertaken by the sectoral agencies and the regional states.
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* This chapter was prepared by George Sarpong.
I. INTRODUCTION

_Biosecurity_ involves the management of biological risks in a comprehensive manner to achieve food safety and protect animal and plant life and health. It also addresses the associated environmental risks. The assumption is that all these sectors are inextricably linked and that the similarities in their regulatory frameworks demand a unified and coordinated approach.

As an issue of global importance, the different components of _biosecurity_ have been addressed in several international instruments, which were reviewed in Chapter 2. This chapter examines Ghanaian legislation in the light of these international instruments and addresses gaps/weaknesses in the Ghanaian regime against the backdrop of the relevant international dictates. The discussion proceeds along two broad but inter-related headings: (1) Ghana and the regime of the World Trade Organization (WTO) Agreement on the Application of Sanitary and Phytosanitary (SPS) Measures, and (2) the Convention on Biological Diversity (CBD), CBD-related instruments and Ghana. This chapter also examines the relevant constitutional and institutional bases for _biosecurity_ in Ghana. It draws conclusions and proffers a way forward for the _biosecurity_ regime in the country.

II. THE GHANAIAN REGIME ON BIOSECURITY

2.1. Ghana and the SPS regime

Various pieces of legislation exist on the statute books to address food safety, animal and plant health in Ghana.

2.1.1. Food safety

In the area of food control and safety, the 1992 Constitution and certain pieces of food safety legislation have a bearing on the subject. The Constitution, the fundamental law of the land, does not expressly address food safety. Some of its provisions, however, are relevant to the subject. Article 36(2)(b) of the Directive Principles of State Policy enjoins the state to take necessary steps to establish a sound and healthy economy whose underlying principles shall include affording ample opportunity for individual initiative and creativity in economic activities and fostering an enabling environment for a pronounced role of the private sector in the economy.
Furthermore, article 36(10) enjoins the state to safeguard the health, safety and welfare of all persons in employment, and to establish the basis of the full deployment of the creative potential of all workers. This provision, which is of significant relevance within the context of health and safety at work, is also relevant for present purposes. A law passed with the object of strengthening public health standards among street food vendors, for example, would be perfectly well in accord with this constitutional injunction since many a "person in employment" relies on street foods as a basis for sustenance.

In addition to the constitutional provisions noted above, the following pieces of legislation govern foods in Ghana:

- Food and Drugs Law, 1992 (PNDCL 305B); and
- Food and Drugs (Amendment) Act, 1996 (Act 523).

The Food and Drugs Board (FDB) has also proposed for adoption draft Food and Drugs Regulations, 2000. These are yet to be passed by Parliament but are being used by the FDB in its operations.

The Food and Drugs Law (FDL) was enacted to control the manufacture, importation, exportation, distribution, use and advertisement of foods, drugs, cosmetics, chemical substances and medical devices. Accordingly, the FDL contains prohibitions against the sale of unwholesome, poisonous and adulterated foods and also prescribes standards for foods.

The definition of "food" is rather wide under the law. Section 51 states that it includes "any article manufactured, sold or represented for use as food or drink for human consumption, chewing gum, water and any ingredient of such food drink, chewing gum or water". This definition is in consonance with accepted practice, as for example obtains under United Kingdom legislation.

The FDL has been amended by the Food and Drugs (Amendment) Act, 1996 (Act 523), whose purpose is to provide for the fortification of salt to alleviate nutritional deficiencies and to bring the provisions of the FDL in conformity with the Constitution. Act 523 expands on the definition of "food" to include "salt and any article manufactured, sold or represented for use as food or drink for human or animal consumption, chewing gum, water and any ingredient of the food, drink, chewing gum or water".
Section 27 of the FDL establishes an administrative authority, the Food and Drugs Board (FDB), under the control and supervision of the ministry responsible for health. The composition of the board is wide-ranging, drawn from relevant departments and agencies of state and the private sector. The functions of the FDB as set out in Section 28 include advising the ministry on all matters relating to the administration and implementation of the FDL.

A major defect of the FDL from the standpoint of Biosecurity is that the FDL is bereft of contents as regards the international standards that should guide the FDB in the discharge of its duties. Schedule I of the law makes reference to the publications of certain international bodies, but the list does not include those of the Codex Alimentarius Commission. An amendment of the FDL would bring it in line with international practice on the subject.

Authorized officers of the FDB have wide enforcement powers under the FDL for purposes of entering premises, opening and examining food receptacles and books and seizing and destroying unwholesome, poisonous or adulterated foods. Obstruction of an authorized officer or contravention of the provisions of the law is an offence punishable, upon summary conviction, by a fine or imprisonment. Individuals, corporations and partnerships are liable to sanctions under the law. Section 35 of the FDL mandates an authorized officer at any reasonable hour to inspect any animal intended for slaughter and to seize and inspect any meat which he or she considers unfit for human consumption.

2.1.2. Animal health

With regard to animal health, few statutes exist, one dating from the colonial era. Of relevance for present purposes are the:

- Animals (Control of Importation) Ordinance (Chapter 247); and

A draft Meat Inspection Law, 1999, has also been proposed by the sector ministry although it is yet to be passed as an Act of Parliament.

The Disease of Animals Act gives the minister responsible for agriculture power to adopt measures to curb the outbreak of animal diseases. Veterinary

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1 For a discussion of the Codex Alimentarius, see Chapter 2, Part V.
officers have powers of inspection over animals. Thus by implication the
draft law seeks to repeal Section 35 of the FDL. The powers exercisable by
veterinary officers under Act 83, especially in the event of outbreak of diseases,
are aimed at the control and avoidance of the spread of animal diseases, and
hence safeguard human health since animals are sources of food.

Contravention of the provisions of the act constitutes an offence, and upon
conviction, attracts imprisonment for a term not exceeding six months or to
a fine not exceeding 50 pounds or to both. The applicable financial sanctions
are, however, outdated and need to be revised. There is also the need to refer
to and incorporate the OIE standards which are the international norms or
benchmarks on animal health and animal diseases.2

The draft Meat Inspection Law was prepared as far back as 1999 to address
the subject of meat inspection. An accompanying memorandum to the law
provides justification for its enactment:

- the need to enact laws and regulations to protect consumers from
  chemical and biological agents harmful to human health;
- the need to entrust the responsibility for meat laws to a government
department responsible for promotion of animal and human health
  – Veterinary Services is seen as best suited in this regard;
- the need to provide legal backing to veterinary personnel of the
  Ministry of Food and Agriculture (MOFA) to effectively carry out
  the meat inspection function;
- the need to ensure compliance of Ghanaian legislation with
  international standards and practices.

The draft legislation is in four parts: Part I deals with the controlling
authority, appointment, qualifications and functions of veterinary inspectors.
Part II deals with inspections, labelling and enforcement of the law. Part III
deals with importation and exportation of meat and other animal products,
and offences. Part IV addresses regulations and interpretation of the law.

Under Part I, the Veterinary Service Directorate (VSD) of MOFA is
designated as the controlling authority responsible for the control of meat
hygiene, including meat inspection, as well as all decisions relating to human
health and animal health at admission of slaughter animals to the abattoir or

2 The OIE, or World Organization for Animal Health, is discussed in Chapter 2, Part IV.
slaughterhouse, and ante-mortem and post-mortem inspections. These provisions are obviously designed to put to rest any controversy as regards the competent authority on the subject. The OIE has indicated that veterinary services departments should exercise responsibility for meat inspection. However, it should be kept in mind that the VSD alone cannot perform this function efficiently and effectively without the assistance or collaboration of officials from other ministries, departments and agencies.

Enforcement

Under the existing regime, both the FDB and the district/metropolitan assemblies have statutory functions in meat inspection. The draft law seeks to divest these bodies of these functions and vest same in the VSD. The justification for this proposal as contained in the memorandum is that VSD "is adequately equipped to undertake the task ... It has qualified staff that can detect diseases in animal[s], recognize normal and abnormal tissues in organs. There are over 700 … technical and professional staff spread throughout the country who can competently and completely take over meat inspection".

The draft in this part also makes provision for the appointment and qualifications of veterinary inspectors. These include a qualified and registered veterinarian and any other veterinary personnel appointed as inspector pursuant to the law. These are accorded wide inspection and enforcement powers under the act. Because the VSD does not have sufficient staffing, a convenient arrangement on the matter could be the sharing of responsibilities between officials of the VSD, public health officers and the FDB.

Part III of the draft law prohibits the importation and exportation of meat and meat products unless certain conditions are met. Meat and meat products cannot be imported unless they bear a certificate testifying that the products satisfy the requirements of standards prescribed by the appropriate authority in Ghana. In the case of exports, there is a similar prohibition unless the products are certified by the Director of Veterinary Services as meeting the prescribed standards set by the Ghana Standards Board.

The Animals (Control of Importation) Ordinance (Chapter 247), a colonial statute which is still in force, bans the importation of animals into the country unless certified by a veterinary authority that they are free from diseases. The ordinance is outdated and could be repealed since the draft
Meat Inspection Law deals with the subject of importation and exportation of meat, meat products and animals.

Even though standards for inspections, importation and exportation of meat products have been prescribed by the appropriate international bodies, there is no express reference to the OIE in the legislation. A reference to these standards could be made in the parent enactment, or regulations could be passed incorporating these standards. Another gap that needs to be addressed is the absence of legislation on animal feeds and veterinary drugs. These could be provided for by way of regulations made pursuant to the proposed law (as provision is made under the draft law for regulations) or by separate legislation addressing each of these topics.

A revised draft meat inspection law prepared in 2004 is an improvement on the 1999 draft. It provides for the appointment of inspectors, meat inspections, importation and exportation of meat and miscellaneous matters including advertising, seizures and regulations. The minister is given the power to make regulations on a wide range of matters including inspections; the registration of establishments and the licensing of operators; fees; and standards for imported meat products. However, the draft still suffers from a failure to refer to international standards as the basis for standard setting and enforcement.

2.1.3. Plant protection

The major piece of legislation governing plants and plant protection is the Prevention and Control of Pests and Diseases of Plants Act (1965). The main objective of the act is to consolidate, with amendments, the legal framework relating to the prevention and treatment of plant pests and that relating to plant quarantine. It has three parts: Part I on prevention and treatment of plant pests; Part II on plant quarantine; and Part III, with miscellaneous provisions.

Part I (sects. 1–8) places a ban on the keeping, selling, offering for sale or barter or distribution of any plant infested or infected with such pests as may be prescribed. Infested or infected plants may not be removed from any land except for the purpose of inspection by an inspector; destruction; or preservation of the produce of such plants for subsequent manufacture for sale or for seed. Provision is made under the act for the making of regulations by the minister on the prevention and eradication of plant pests and other relevant matters.
Part II of the Act (sects. 9–12) prohibits any person, except with a permit, from importing any plants, plant products, plant pests, soil, manure, grass, packing materials or any other material liable to harbour dangerous plant pests. Provision is made for regulations prescribing the conditions for the grant of a permit, examination of items imported under the permit and the prohibition and restriction of importation of such items. Inspectors have powers to seize items being imported in contravention of the act and to dispose of them.

Part III of the Act (sects. 13–17) makes it an offence punishable on conviction to a fine not exceeding 100 cedis or to a term not exceeding three months to contravene a direction, requirement, condition or prohibition imposed under the act. Provision is also made for compensation to occupiers of land or importers who may suffer pecuniary losses as a result of measures taken under the act.

Other relevant legislation is the Seed Inspection and Certification Decree, 1972, and the Pesticides Control and Management Act, 1996. Together with Act 307, they provide the legal and institutional bases for the coordination and regulation of plant protection activities in Ghana.

Enforcement

Within MOFA, the Plant Protection and Regulatory Services Department (PPRSD) is responsible for plant protection, plant quarantine, seed quality control and pest management. Officers of the PPRSD carry out phytosanitary inspections at all border points including the sea ports and the international airport. The Customs, Excise and Preventive Service (CEPS) officers, also stationed at the borders, are required to inform the plant quarantine officers if they notice any plant materials in imported shipments or baggage which they (CEPS) have inspected in the first instance. With regard to risk assessment, the PPRSD undertakes this function through the employment of a core of trained risk assessors drawn from MOFA, research institutions and the universities.

Proposal for reform of the law

The legal regime for plant protection is outdated, and the legislation has several drawbacks, and does not measure up to the standards of the new
revised text of the International Plant Protection Convention (IPPC). The shortcomings include absence of provisions on risk analysis and exportation of plants, lack of adequate provisions on the duties of plant quarantine officers and on co-ordination among the various institutions involved in plant protection in the country as well as low financial penalties for violations.

Under the FAO Technical Co-operation Programme, a project, "Strengthening Plant Quarantine Capabilities: Republic of Ghana", was initiated in 1996. The main output of the project, a draft Plant Quarantine Act, was presented to the government for consideration and adoption. The draft act seeks to repeal the 1965 Plant Protection Act and replace it with a modern Plant Quarantine Act that will prevent the introduction and spread of plant pests in Ghana.

The 1996 draft act has itself been overtaken by the recent coming into force of the new revised text of the IPPC and needs to be updated to incorporate provisions and concepts of the new text, including an enhanced role for the national plant protection organization (PPRSD) in Biosecurity, such as through collaboration with the Ministry of Environment on living modified organisms and ecological risks.

2.2. Ghana, the CBD and CBD-related instruments

Ghana has not enacted specific legislation to domesticate the Convention on Biological Diversity (CBD). However, several pieces of legislation along sectoral lines – fisheries, forestry, game and wildlife – exist on the statute books to address the subject. In general these are designed to ensure the conservation and sustainable utilization of these resources. For present purposes, two pieces of legislation, one on wetlands and a draft Biosafety Bill, are of direct relevance.

On wetlands, the Wetland Management (Ramsar Sites) Regulations, 1999 (LI 1659), have been passed to domesticate the Ramsar Convention. Under these regulations, six Ramsar sites have been designated, namely, Densu Delta, Keta Lagoon Complex, Muni-Pomadze, Owabi Wildlife Sanctuary, Sakumo and Songor. The regulations make provision for declaration of closed seasons and the designation of core areas, authorized activities, proscribed activities

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3 For a discussion of the IPPC, see Chapter 2, Part III.
4 For a discussion of the CBD, see Chapter 2, Part VI.
and restricted activities in the designated areas. The regulations also makes provision for offences for contravention of its provisions and prescribes penalties in the form of fines, imprisonment or both, depending on the gravity of the infraction. The Wild Life Division of the Ministry of Lands, Forestry and Mines is the responsible implementing authority for the regulations.

The draft Biosafety Act, 2004, is designed to domesticate the Cartagena Protocol on Biosafety.\(^5\) The draft act is the outcome of consultations and surveys involving the major stakeholders in the public and private sectors as well as the general public. Its objectives are to ensure, in accordance with the precautionary principle, an adequate level of protection in the field of safe transfer, handling and use of genetically modified organisms (GMOs) resulting from modern biotechnology that may have an adverse effect on the environment; and to establish a transparent and predictable process to review and make decisions on GMOs and related activities.

The draft’s 44 sections and four schedules address all the requirements for the implementation of the protocol. Among other things, the draft act creates an institutional framework, the National Biosafety Authority (NBA), a corporate entity with perpetual succession, with representation drawn from both the public and private sectors. The functions of the NBA are to:

- receive, respond to and to make decisions on applications under and in conformity with the act;
- establish administrative mechanisms to ensure the appropriate handling and storage of documents and data in connection with the processing of applications and any other matters covered by the act; and
- promote public awareness, participation and education concerning the activities of the NBA under the act.

The draft act creates a regulatory regime with accompanying regulations to address the handling of permits, monitoring and enforcement, approvals and appeals, public participation and information and finance. Decisions on GMOs are to be based on science – risk assessment – and the procedure is set out in the fourth schedule to the act to ensure transparency and predictability.

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\(^5\) For a discussion of the protocol, see Chapter 2, Part VII.
Taking cognizance of the fact that biosafety is a multi-institutional activity that cuts across several sectors, the draft act does not re-invent the wheel: it employs the services of the existing regulatory agencies in the Technical Advisory Committee (TAC). The functions of the TAC are to act as the national advisory committee on matters concerning or related to the genetic modification of organisms, carry out risk assessment of applications at the request of the NBA and advise, upon request or on its own accord, the minister, the NBA, the ministries and appropriate bodies on matters concerning the genetic modification of organisms.

The inclusion of the regulatory agencies in the TAC should help ensure the much-needed coordination and cross-sectoral management required in such a multi-institutional endeavour. It should also help to address the overlaps, conflicts and gaps in mandates of the various regulatory agencies that now operate within the TAC. This kind of arrangement has much to commend itself especially for developing countries that lack the requisite capacity to create new institutions to address Biosecurity. Indeed an examination of the current institutional arrangements for Biosecurity reveals overlaps and conflicts in the mandates of the various institutions whose functions have a bearing on the subject, a situation that impedes its smooth operation.

2.3. The institutional basis for Biosecurity in Ghana

Biosecurity issues are not the responsibility of one ministry, department or agency (MDA) of state. Indeed, several MDAs have responsibility for, or are engaged in, activities in this area. These include:

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<td>- FDB</td>
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<tr>
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<tr>
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Ministry of Finance  
- Quality Control Division of Cocoa Marketing Board  
- Cocoa Research Institute

Ministry of Trade, Industry, Private Sector and Presidential Special Initiatives  
- GSB  
- CEPS

Ministry of Tourism and Diasporean Relations  
- Ghana Tourist Control Board

Ministry of Education
Ministry of Fisheries
Ministry of Foreign Affairs
Ministry of Justice
Ministry of National Security
Ministry of Water Resources, Works and Housing

Under the Constitution, the executive power of state is vested in the President (art. 58). The MDAs are part of the executive arm of government. Departments, agencies and institutions such as EPA and CEPS are the creatures of statutes, and their functions or mandates are provided for in the enactments that established them. The Constitution also makes provision for the establishment of the:

- Lands Commission;
- Minerals Commission;
- Forestry Commission;
- Fisheries Commission; and
- such other commissions as parliament may determine.

These commissions are entrusted with responsibility for the regulation and management of the utilization of the respective natural resources and the coordination of policies in relation to them (arts. 258 and 59). Parliament has since enacted appropriate legislation to establish these commissions and other regulatory bodies. Of these, the most relevant for present purposes are:

- FDB and GSB – responsible for food and related safety matters;
- VSD – responsible for animal health and related safety issues;
- PPRSD – responsible for plant health and related safety matters;
EPA – responsible for environmental regulation, including safety matters;
CEPS – responsible for ports and borders in collaboration with the other agencies; and
district, municipal and metropolitan assemblies which work with the regulatory agencies at the local levels of governance especially in monitoring and enforcement at markets.

These institutions are responsible for the fulfilment of Ghana’s obligations under the WTO SPS regime: the PPRSD is the mandated national plant protection organization; the FDB and GSB are responsible for the Codex Alimentarius and the WTO Agreement on Technical Barriers to Trade, respectively; and the VSD is the responsible agency for the OIE. The overall regulatory/coordinating body, however, is EPA.

**Environmental Protection Agency (EPA)**

EPA was established pursuant to the Environmental Protection Agency (EPA) Act, 1994 (Act 490). Among its functions is the issuance of environmental permits and notices for controlling the volume, types, constituents and effects of waste discharges; prescription of standards and guidelines relating to pollution of water; and enforcement of environmental standards including ensuring compliance with established environmental impact assessment (EIA) procedures in the planning and execution of projects.

In pursuance of these functions, EPA officers embark on inspections in hotels and other industrial establishments to ensure maintenance of prescribed effluent standards and standards of hygiene generally. An issue that has serious implications for food safety is the contamination of food, in particular pesticides residues in food. The subject is addressed by the Hazardous Chemicals Committee (HCC) established pursuant to Section 10 of the EPA Act. The duties of the HCC are to:

- monitor the use of hazardous chemicals by collecting information on their importation, exportation, manufacture, distribution, sale, use and disposal;
- advise the board and the executive director of EPA on the regulation and management of hazardous chemicals;
- perform such other functions relating to such chemicals as the board or the executive director may determine.
The subject of pesticides is further regulated under the Pesticides Management and Control Act, 1996 (Act 528). Under Section 17, no person shall import, export, manufacture, distribute, advertise or sell any pesticides except in accordance with a licence issued under the act. The licensing regime is under the overall management and control of EPA.

A related problem is that of certification of agricultural exports as being free from pesticide residues for purposes of gaining access to foreign markets. This is an issue of extreme importance in view of the fact that in recent times, Ghanaian agricultural produce has been rejected in Europe on account of high residue levels. To date, no regulations have been enacted pursuant to Act 528 either on pesticide residues in food or on certification of agricultural produce as being free from residues.

With regard to cocoa, however, which is Ghana’s major export crop, the Cocoa Research Institute of Ghana (CRIG) has the requisite facilities for vetting the effectiveness, toxicity and possible residues of pesticides that has been applied. CRIG works in close collaboration with EPA on the subject. In the event of a disagreement between CRIG and EPA with respect to residues on cocoa, an application cannot be granted. The GSB and the Ghana Atomic Energy Commission may also veto the registration of a pesticide. As in the case of other agricultural produce, there is the need to embody these standards in subsidiary legislation pursuant to Act 528.

2.4. Conflicts, gaps and overlaps

The current institutional arrangements for Biosecurity are bedevilled with gaps, overlaps and conflicts in the mandates of the various institutions whose functions have a bearing on Biosecurity, as illustrated below.

At the borders, PPRSD officials carry out inspections of all imported goods thought to consist of plants and plant materials. The PPRSD has 44 entry points around the country where their personnel are located. CEPS officers, also stationed at the borders, are required to inform PPRSD if they notice any plant materials in imported shipments or baggage which they (CEPS) have inspected in the first instance. The effectiveness of this collaboration depends on the particular border post in question. At Kotoka International Airport, the two services work well together. At Tema Port (the major sea port) by contrast, large shipments of plant materials are released into the country without the PPRSD inspecting the shipments or even being informed that the
shipments have arrived. Another problem facing both CEPS and the PPRSD is that there are several unapproved entry points into the country.

As part of the efforts towards improving the management of the country’s borders, the Government of Ghana has decided to enhance the capacity of the Ghana Immigration Service (GIS) to perform more efficiently and effectively its responsibilities with respect to the entry and exit of persons to and from the country. This will enable GIS to deal with cross-border and travel-related crimes with a high degree of accuracy and efficiency. As part of the measures, a Border Patrol Unit has been established to intercept any illegal entries into the country. The idea is to manage immigration as an integral part of the national development agenda. These measures require a much closer collaboration between GIS and the other regulatory agencies operating at the borders.

The Ghana Tourist Control Board (GTCB) established by the Tourist Board Decree, 1973 (NRCD 224), as amended by the Ghana Tourist Control Board (Amendment) Decree, 1977 (SMCD 80), has among its functions the registration, classification, licensing and control of standards in hotel accommodation and catering enterprises. In pursuance of this function, officials of the GTCB conduct periodic inspections of hotels and catering businesses to ensure compliance with its standards. There appears to be little co-ordination/co-operation between officials of the GTCB, public health officials and FDB officials. In view of the implications of these inspections for food safety, there is the need for close collaboration and overall coordination between the GTCB and these regulatory agencies.

The meat inspection function is a source of conflict/overlap between the veterinary services and the public health officers (PHOs) of the metropolitan and district assemblies. The meat inspection function has been exercised by PHOs by virtue of previous and current legislation on local government. Unlike these officials whose mandates are expressly provided for in LI 1615, no such specific mandate is accorded to veterinary officers in the area of meat inspections. As noted, both the FDB and the district/metropolitan assemblies (local government institutions) have statutory functions in meat inspection. The draft law seeks to divest these bodies of these functions and vest same in the VSD, but makes provision for the inclusion of inspectors from the district assemblies in the meat inspection function.
The Ghana Standards Board (GSB) was established by the Standards Decree, 1967 (NLCD 199), as a body corporate with perpetual succession. NLCD 199 was repealed by the Standards Decree, 1973 (NRCD 173). NRCD 173 continued the existence of the GSB as a body corporate with perpetual succession. Section 3 of NRCD 173 provides for a wide range of functions and powers of the GSB including:

- recommending to the ministry responsible for industries to prohibit the sale or manufacture of goods in the national interest as well as in the interest of public health and safety;
- maintaining the necessary machinery to ensure that goods prepared and manufactured for export are distinctly marked for export only;
- providing for the issuance of a certificate to the effect that goods comply with known requirements or standards in the country to which they are to be consigned, before the export of such goods is permitted; and
- recommending to the ministry responsible for trade to prohibit the importation into Ghana for the purposes of sale, use or human consumption, any goods unless the same have been certified by the GSB as complying with standards set up by the GSB.

Section 9 of the decree gave the board of the GSB power to make, alter or rescind rules on a wide range of matters, including providing for the amendment or revocation of any standard. NRCD 173 was subsequently amended by AFRCD 44, with the addition of two specific functions on food to be exercised by the GSB, namely: prohibiting the sale or manufacture of foods in the national interest; and prohibiting the importation into Ghana of foods which have not been certified by the board as complying with its standards.

From the above it is apparent that the GSB, like the FDB, has statutory functions in the area of sale, manufacture, exportation and importation of foods. It is these provisions which have become a source of overlap/conflict between the GSB and the FDB, the latter of which was established in 1992 to control the manufacture, importation, exportation, distribution, use and advertisement of foods, drugs, cosmetics, chemical substances and medical devices.
Apart from the overlaps, the existing laws on standards are also deficient in matters such as low fines for offences and low licence fees. Furthermore, even though in practice the standards sought to be implemented under the law are based on international standards such as the Codex Alimentarius, there is no explicit reference to any of these standards in any of the existing legislation.

A draft standards bill has been proposed to address the foregoing and other defects in the existing legal regime for standards. The draft bill establishes a National Standards Authority (NSA) as a body corporate with perpetual succession and enhanced powers of acquisition of land.

The draft bill re-enacts the provisions on the functions of the GSB under NRCD 173 as amended by AFRCD 44. The specifications for standards to be prescribed under the law include "international or other overseas specifications." In view of the fact that the WTO has prescribed the TBT Agreement as the source of international norms/benchmarks for goods other than those prescribed by the Codex Alimentarius, explicit reference to WTO standards would be appropriate.

A thorny issue that the draft bill does not satisfactorily address is the competence/functions of the NSA in the area of foods vis-à-vis the FDB. Under Section 3(2)(c) of the draft bill, the NSA is authorized to prohibit, in the national interest, the sale or manufacture of any kind of goods. It also has power to prohibit the importation of goods that have not been certified by the authority as complying with its standards. The definition of "goods" is wide enough to encompass foods. Hence the draft bill in its present form conflicts with the mandate of the FDB which, as noted, has been established to control the manufacture, importation, exportation, distribution and use of foods.

The draft bill in its fourth schedule repeals the Standards Decree, 1973. It, however, continues in force any statutory instruments in force under the prior decree. The import of this provision is that certain regulations affecting foods would continue to remain in force. This calls for amendments to be made to the regulations, to remove all references to food, otherwise there will continue to be overlaps with the FDB. Further, in line with GSB's policy of developing voluntary standards in conformity with WTO guidelines, there is the need for regulations on certification of goods when the goods have been prescribed to have health and safety implications for consumers.
III. THE GHANAIAN BIOSECURITY FRAMEWORK: CONCLUSIONS AND THE WAY FORWARD

Biosecurity has assumed great importance in the twenty-first century owing to rapid technological advances and increased international trade. The international multilateral treaty regimes, including the WTO and the CBD, have prescribed or endorsed rules to address the subject. This chapter has assessed the existing Ghanaian constitutional, institutional and legislative frameworks in light of the relevant international standards. The discussion on the legislative framework was conducted along two broad but interrelated headings: Ghana and the SPS regime; and the CBD and CBD-related instruments and Ghana. Several results emerge from the study.

In relation to the SPS regime, several pieces of legislation exist which address food safety, plant and animal health. They, however, fall short of the standards prescribed by the relevant international bodies – Codex, the IPPC and the OIE. With regard to the CBD and CBD-related instruments, no single piece of legislation addresses the subject; there are several pieces of sectoral legislation on fisheries, forestry, game and wildlife. Indeed, the challenge faced by the government is how to implement these standards effectively given the limited resources available. However, specific legislation has been enacted to domesticate the Ramsar Convention on Wetlands, and comprehensive draft legislation has been prepared for the implementation of the Cartagena Protocol on Biosafety.

Some attempts have been made to revise existing legislation on food safety, plant and animal health. The outcomes have not been satisfactory: the proposed drafts do not measure up to the international norms prescribed by the SPS regime or they have been overtaken by subsequent developments on the subject. The draft Biosafety Act, 2004, designed to domesticate the Cartagena Protocol, however, is comprehensive and meets the objects and requirements of the protocol. Its institutional arrangement, involving the employment of all the existing regulatory agencies under the aegis of the NBA, points to a way of resolving the conflicts, gaps and overlaps in the Ghanaian regime on Biosecurity.

Biosecurity is not the responsibility of one agency of state; it involves several ministries, departments and agencies. The regulatory framework for Biosecurity is multi-sectoral in nature, but without an overall coordinating
body in Ghana. As a result, the regime is bedevilled with overlaps, gaps and conflicts among and between the regulatory agencies.

The challenge faced by Ghana is thus not only one of enacting legislation consistent with the prescribed international norms, but also ensuring a coordination of the regime so as to eliminate the various gaps, overlaps and conflicts. Indeed the need to implement WTO standards apart from the health implications for the population has serious implications for trade between Ghana and the North as well.

Even though developing countries now have preferential treatment in terms of lower or zero tariffs and non-tariff barriers for their products, similar concessions cannot be gained over sanitary and phytosanitary standards which are regarded as highly sensitive in view of their health implications. Once the standards are in place, the only options are either to meet those standards and export accordingly or drop the idea of exporting altogether. At the end of the day, no consumer would buy any agricultural product that does not conform to prevailing standards. Indeed, no such concessions can be gained even under the preferential concessionary arrangements under the ACP (African, Caribbean and Pacific Group of States)/European Union framework.

Ghana should thus strive to implement WTO rules in domestic legislation, in large part because they constitute treaty obligations assumed under the WTO. There are, however, constraints, including limited technical, human and financial resources. These constraints could be addressed by a programme of capacity building, through international and donor collaboration with organizations such as the WTO, the United Nations Environment Programme, FAO and the World Health Organization.

In the development of an appropriate legal framework for Biosecurity, two institutional options, in the light of the analysis, are worthy of consideration: the creation of a new body at a higher level, for example, under the Presidency, to exercise an oversight and coordinating role over Biosecurity; and the use of existing or proposed coordinating frameworks to address the subject.

The creation of a new body at a high level of state such as the Presidency to ensure coordination of, or to exercise oversight responsibility over, Biosecurity accords the issue proper attention. It will also prevent ministries from vying
with one another for resources and power. The creation of a bureaucracy to deal with the subject, however, is likely to entail significant costs.

The other option would be to enhance the functions of the proposed NBA to deal with Biosecurity as a whole. In this regard, the composition and functions of the NBA, its TAC and inspectorate could be enhanced to include representation from other Biosecurity agencies not represented in the NBA, such as national security and the GIS. Such a measure would be cost-effective. Indeed, the regulatory framework provided for under the draft Biosafety Act provides, with the necessary modification, a means of ensuring the much-needed coordination and cross-sectoral management required in such a multi-institutional activity as Biosecurity.

This kind of arrangement has much to commend itself for application by a developing country like Ghana that faces constraints in the creation of a new institution to address Biosecurity. Since the legislation is still in draft there will not be the requirement for any amendment to effect the proposed changes. They could be incorporated in the draft for appropriate parliamentary action.
INDIA COUNTRY STUDY

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* This chapter was prepared by Roopa Madhav and Adil Hasan Khan.
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I. REFERENCE TO BIOSECURITY IN INDIA

In recent years, reference has been made in two policy documents to the need to bring about comprehensive legislation dealing with Biosecurity in India. The first such document was the May 2004 Report of the Task Force on Agricultural Biotechnology.1 The report advised the Government of India to prepare a Biosecurity Compact in order to deal with the following issues:

- invasive alien species;
- sanitary and phytosanitary measures to avoid mycotoxins, salmonella and other forms of infection in food;
- food, environment and biosafety relating to genetically modified organisms (GMOs); and
- bio-ethical considerations in research.2

The report recommends setting up a task for the preparation of such a Biosecurity Compact.

Most recently, the Revised Draft National Policy for Farmers, issued in October 2006, includes among its ten major goals strengthening the "Biosecurity of crops, farm animals, fish and forest trees for safeguarding both the work and income security of farmer families, and the health and trade security of the nation".3 The document calls for the creation of the National Agricultural Biosecurity System (NABS), with the following aims:

"Safeguard the income and livelihood security of farmer families, as well as the food, health and trade security of the nation, through effective and integrated surveillance, vigilance, prevention and control mechanisms designed to protect the productivity and safety of crops, farm animals, fishes and forest trees."

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2 Id.
"Enhance national and local level capacity in initiating proactive measures in the areas of monitoring, early warning, education, research, control and international cooperation."

"Introduce an integrated Biosecurity package comprising regulatory measures, education, improved sanitary and phytosanitary measures and social mobilization."

"Organize a coordinated National Agricultural Biosecurity Programme on a hub and spokes model, with effective home and regional quarantine facilities capable of insulating the major agro-ecological and farming systems zones of the country from invasive alien species of pests, pathogens and weeds as well as from the introduction and release of GMOs".4

Biosecurity was also added as an area of cooperation under the US-India Agricultural Knowledge Initiative in June 2006, which aims to address the issue, starting with threat posed to crops by invasive alien species up to averting the release of bio-agents of mass destruction.5

II. BIOSECURITY LAWS IN INDIA

India has a plethora of laws which deal with Biosecurity but it needs to be noted that they do not stem from an understanding of the term. The pieces of legislation have been enacted with differing objectives and public concerns in mind. Though disparate and scattered, these pieces of legislation serve an essential function in specifically addressing the sectoral concerns, and they carry forth the intent contained in the preambles. Likewise, the institutions, though numerous, have been established to serve the purposes of the original enactments.

2.1. Constitution of India

Though there is no specific reference or use of the term Biosecurity in the Constitution of India, a number of its provisions are of relevance to understanding the legal framework dealing with Biosecurity in the country. The Constitution is also the key to understanding how the general legal set-up works.

4 Id.
2.1.1. Directive Principles of State Policy

Part IV of the Constitution contains the Directive Principles of State Policy. Within these, article 47 is relevant and it, among other things, makes it the duty of the state to improve public health. Article 48 is also of relevance as it provides that the state shall endeavour to organize the agricultural and animal husbandry sectors on modern and scientific lines. Article 48A, which was inserted by the 42nd Amendment to the Constitution in 1976, requires the state to "protect the environment and to safeguard the forests and wild life of the country".

2.1.2. Fundamental rights

Part III of the Constitution of India contains the fundamental rights. Among these is the right to life, which is enshrined in article 21, and which has the most relevance for the legal framework for Biosecurity. Since the late 1970s, the Supreme Court, which is the highest court of the country, has progressively widened the scope of the rights granted under this article. This has been achieved by giving an expansive interpretation of the term "life". As a result of judicial interpretation, the right to life has become a sort of repository of various human rights. Some of the pertinent rights thus included are:

- the right to health;
- the right to a healthy environment;
- the right to pollution-free water and air; and
- protection against hazardous industries.

2.1.3. Federal scheme

Since India has a federal Constitution, it necessarily provides for a division of power and functions between the centre and the federal units (states). The Indian federal system leans slightly in favour of the centre while keeping a federal pattern and framework. The Constitution has created three functional areas regarding law-making by the two components of the federal system. These are:

- an exclusive area for the centre called the Union List;
- an exclusive area for the states called the State List; and
• a common or concurrent area in which both the centre and the states may operate simultaneously, though with the centre having overall supremacy, called the Concurrent List.

The relevant article of the Constitution in this regard is article 246, which creates this scheme of division and flexible sharing. The actual lists are provided in the seventh schedule of the Constitution. As far as Biosecurity is concerned the relevant entries are:

List I – Union List
Entry 28. Port quarantine, including hospitals connected therewith.
Entry 51. Establishment of quality standards for goods to be exported out of India or transported from one state to another.

List II- State List
Entry 6. Public health and sanitation; hospitals and dispensaries.
Entry 14. Agriculture, including agricultural education and research, protection against pests and prevention of plant diseases.
Entry 15. Preservation, protection and improvement of stock and prevention of animal diseases; veterinary training and practice.

List III- Concurrent List
Entry 17A. Forests.
Entry 17B. Protection of wild animals and birds.
Entry 18. Adulteration of foodstuffs and other goods.
Entry 29. Prevention of the extension from one state to another of infectious or contagious diseases or pests affecting men, animals or plants.

2.1.4. International law

As per article 253 of the Constitution, the Indian Parliament has been given the power to enact any law to implement the international treaties, conventions or agreements entered into with other countries or even decisions made at any international conference, association or other body. This power is not affected by the subject matter of the legislation. That is, if India becomes a party to any international convention, parliament can enact a law to effectuate its obligations under the same, even if the subject matter of the enactment is specifically one that, according to the lists, falls within a different domain.
However, it must be kept in mind that the parliament’s power to legislate in respect of an international treaty entered into by the state is not unlimited and is limited by other constitutional restrictions, e.g. fundamental rights.

2.2. Food safety

2.2.1. Legislation

Food Safety and Standards Act, 2006

The Food Safety and Standards Act consolidates the laws governing the food sector. The act establishes the Food Safety and Standards Authority of India (FSSAI), which is assisted by a central advisory committee, a scientific committee and several scientific panels. The FSSAI shall lay down science-based standards for food articles and seeks to regulate their manufacture, import, storage, distribution and sale, to ensure availability of safe and wholesome food for human consumption.

The act defines "food" to mean any substance, whether processed, partially processed or unprocessed, which is intended for human consumption and includes primary food, genetically modified or engineered food or food containing such ingredients, infant food, packaged drinking water, alcoholic drink, chewing gum and any substance, including water, used in the food during its manufacture, preparation or treatment (sect. 3(j)).

Section 3(s) states that the "Food Safety Management System" means the adoption of Good Manufacturing Practices (GMP), Good Hygienic Practices (GHP), Hazard Analysis and Critical Control Point (HACCP) and such other practices as may be specified by regulation, for food businesses.

The FSSAI is to be assisted by several scientific panels and a central advisory committee in laying down standards for food safety and in its overall functioning. These standards will include specifications for ingredients, contaminants, pesticide residues, biological hazards and labels. The act empowers State Commissioners of Food Safety and other local-level officials to implement the law.

Every entity in the food sector is required to get a licence or registration from local authorities. Every distributor is required to be able to identify any
food article by its manufacturer, and every seller to identify any food article by its distributor. Any entity in the sector is bound to initiate recall procedures if it finds that the food sold has violated specified standards. The Commissioner of Food Safety (CFS) of each state, through food safety officers (FSOs), enforces the standards.

The act prohibits the use of food additives, processing aids, contaminants, heavy metals, insecticides, pesticides, veterinary drugs residue, antibiotic residues or solvent residues unless they are in accordance with specified regulations. Certain food items such as irradiated food, genetically modified food, organic food, health supplements and proprietary food cannot be manufactured, processed or sold without adhering to specific regulations.

For a specific district, the CFS of each state appoints a Designated Officer (DO), not below the level of Sub-Divisional Officer, whose duties include issuing or cancelling licences, prohibiting sale of food articles that violate specified standards, receiving reports and samples of food articles from FSOs and getting them analysed. The DO also has the power to serve an "improvement notice" on any food operator and suspend his or her licence in case of failure to comply with such a notice. The DO also investigates any complaint made in writing against FSOs. FSOs are appointed for a specified local area and their duties include taking samples of food articles, seizing food articles that are of suspect quality or inspecting any place where food articles are stored or manufactured.

The act has special provisions for food recall procedures. If a food business operator (i.e. anyone owning or carrying out a business relating to food) considers that a food item is not in compliance with the specified standards, he or she has to initiate procedures to withdraw the food in question and inform the competent authorities.

The act provides for a graded penalty structure where the punishment depends on the severity of the violation. Offences such as manufacturing, selling, storing or importing sub-standard or misbranded food could incur a fine. Offences such as manufacturing, distributing, selling or importing unsafe food causing injury are punishable with imprisonment.
The Prevention of Food Adulteration (PFA) Act was enacted with the objective of assuring the quality and safety of food as well as encouraging fair trade practices. In effect, the statute sought to protect the consumer from the supply of adulterated food by specifying food safety and quality standards for consumer protection. The state governments and the union territories are responsible for monitoring and implementation of the provisions of the PFA Act and Rules.

According to the rules, no person shall manufacture, sell, store or distribute adulterated or misbranded food products not conforming to the prescribed standards. These standards apply to imported food as well as food domestically produced.

The institutional set-up under the PFA Act includes local food inspectors and public analysts, both at the municipal and state levels, their laboratory facilities, the four central food laboratories designated under the PFA Act and the central PFA Division under the Ministry of Health and Family Welfare (MOHFW). The Central PFA Division is also designated the National Codex Contact Point for India.6

The PFA Act provides for the inspection and certification of imported food. It prohibits the import of food which is adulterated, misbranded or which contravenes the provisions of the PFA Act or Rules. The important provisions which are required to be followed essentially while importing/clearing the food products are:

- authorized officers check imported food products;
- the custom collector checks imported food products; and
- authorized officers, on suspicion, may detain any imported food product and send the samples to the Central Food Laboratory for analysis.

MOHFW has prescribed maximum tolerance limits for pesticides and heavy metals in food products under the PFA Rules. MOHFW has also notified draft rules to amend the PFA Rules to regulate the sale and import of food.

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genetically modified or genetically engineered organisms obtained through modern biotechnology and to ensure mandatory labelling of all such products. The purpose is to provide correct information to consumers about the nature of food they purchase for consumption.

*Essential Commodities Act, 1955*

The Essential Commodities Act has been enacted to protect the interests of the general public through the control of the production, supply and distribution of and the trade and commerce in certain commodities. Section 3 of the act empowers the central government to issue control orders for regulating production, distribution, quality, movement and licensing pertaining to essential commodities. Similarly, exercising the powers delegated under the act, the state governments have issued a number of control orders to regulate various aspects of trading in essential commodities such as food grains, edible oils, pulses, kerosene, sugar, etc.

*Other orders*

Several orders were issued under Section 3 of the Essential Commodities Act addressing registration of manufacturers, hygiene in production, labelling and other requirements for specific foods. These include the Vegetable Oil Products (Regulation) Order, 1998, the Milk and Milk Products Order, 1992, the Meat Food Products Order, 1973 and the Fruit Products Order, 1955.

*Export (Quality Control and Inspections) Act, 1963*

The Export Act provides for the sound development of the export trade of India through quality control and inspection. It establishes the Export Inspection Council of India (EICI), which shall, *inter alia*, advise the central government regarding measures for the enforcement of quality control and inspection in relation to commodities intended for export.

Section 6 empowers the central government to (a) notify commodities that shall be subject to quality control or inspection; (b) specify the type of quality control or inspection to be applied to a notified commodity; (c) establish, adopt or recognize one or more standard specifications for a notified commodity; (d) prohibit the export of notified commodities that do not satisfy the quality control or inspection.
The Bureau of Indian Standards (BIS) is a statutory autonomous body set up by this enactment. It comprises members representing industry, consumer organizations, scientific and research institutes, technical institutions, central ministries, state governments and members of parliament.

The BIS provides for quality certifications. It has two kinds of certification schemes: (a) product certification; and (b) management systems certification. The product certification scheme has the primary objective of ensuring quality, safety and dependability for consumers. The scheme, although essentially voluntary, has been made mandatory for certain products such as drinking water, food colours and additives.

The management systems certification (MSC) activity of the BIS consists of a series of activities aimed at assessing the capability of an organization’s management systems such as:

- Food Hygiene – Hazard Analysis and Critical Control Point System – IS 15000: 1998; and
- the combination of two or more systems (integrated management systems).

The MSC activity provides third-party certification to organizations. The Indian Standard on Food Hygiene is technically equivalent to the Codex document on the subject (Codex ALINORM 97/13A).

The Ministry of Commerce and Industry (MOCI) has designated BIS as the enquiry point under the World Trade Organization (WTO) Agreement on Technical Barriers to Trade (TBT Agreement). According to the TBT Agreement, the Enquiry Point issues notifications on proposed technical regulations and certification systems in India to the WTO in Geneva.

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7 See Chapter 2, Section 2.2 for a full description of the TBT Agreement
2.2.2. Institutions

In India, international standards, guidelines, and recommendations are increasingly used to guide domestic as well as international trade. (a) The Directorate General of Health Services (DGHS) in the MOHFW is working to integrate Codex standards into food laws as much as possible. (b) The EICI, the official certification body for exports, is developing standards for exports based mainly on Codex, but it also takes into account that an importing country may impose stiffer requirements. (c) The Codex HACCP and food hygiene standards have been adopted by the BIS. (d) As seen earlier, inspection and certification in India have a regulatory basis under the Export Act of 1963.

The main system of inspection and certification being followed by the EIC in the food sector is food safety management systems-based certification (FSMSC). The FSMSC is aligned with international standards on GMP, GHP and HACCP.

In addition to certifying food products in compulsory areas, the EIC also certifies other products for exports with a focus on the food sector. With the concept of equivalence having been recognized in the WTO Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement) as well as being encouraged at the international level by the Codex Alimentarius Commission, the EIC is emphasizing developing equivalence agreements on conformity assessment with its major trade partners.

The processed food exports from the country are handled by two apex-level agencies, namely the Agricultural and Processed Food Export Development Authority and Marine Products Export Development Authority. The Ministry of Food Processing Industries (MOFPI) is the nodal central government entity proactively involved with the food processing industry in regard to macro policy issues and planning for the sector.

MOFPI is in charge of the implementation of various food safety and quality concerns codified in numerous acts and other government measures. For example, the Fruit Products Order, 1955, promulgated under Section 3 of the Essential Commodities Act, prescribes minimum norms for sanitary and hygienic conditions of manufacturing premises and also lays down product

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8 See Chapter 2, Section 2.1 for a full description of the SPS Agreement.
standards. It is closely associated with the Codex Contact Point in the country, namely the Directorate General of Health Services.

With regard to genetically modified (GM) food, several central ministries and departments are involved in India’s programme of food quality and safety and hence each one of them has a role to play in the activities related to GM foods in India. These include:

- the Ministry of Environment and Forests. This ministry holds the Secretariat of the Genetic Engineering Approval Committee, the apex body that gives approval for manufacture, sale, import and export of all genetically modified organisms (GMOs) and products thereof, including foodstuffs and additives using GMOs or cells;
- the Department of Health in the MOHFW. This department is responsible for implementation of the PFA Act under which the quality and safety of food is regulated;
- the Indian Council of Medical Research (ICMR). This is the apex body in India for the formulation, coordination and promotion of biomedical research under the MOHFW. ICMR acts as an advisory body for MOHFW on various issues including GM foods;
- the Ministry of Agriculture. This ministry comprises three departments, namely the Department of Agriculture and Cooperation, Department of Agricultural Research and Education/Indian Council of Agricultural Research (ICAR) and Department of Animal Husbandry and Dairying;
- MOFPI. This ministry supports the active participation of industry in the laying down of food standards as well as their harmonization with international standards. This ministry is also the licensing authority for processed fruits and vegetable industries; and
- MOCI. This ministry formulates the export policy of the country.

The Central Committee of Food Safety, a legal body under the PFA Act, the Central Fruit Products Advisory Committee and the concerned apex export promotion institutions under the MOCI regularly interact to update and amend existing domestic food laws.

As laid out in the transparency clause (art. 7) and further elaborated in Annex B of the SPS Agreement, the Trade Policy Division (TPD) of MOCI has been designated as the national notification authority (NNA) for the country. The NNA coordinates with different concerned ministries and departments for appointment of enquiry points.
Imported food is inspected at the ports of entry by personnel of the Collectorate of Customs. The Government of India through its various departments – Health, Revenue, Commerce and the Directorate General of Foreign Trade – has initiated several steps to streamline the checking of imported food. As noted earlier, the EICI is the official government inspection body certifying food products for exports.

2.2.3. Evaluation

Within the Indian context, the food safety legislative instruments are presently disparate, with several subordinate rules, regulations and orders having been enacted to deal with contingencies as and when they arose. The operative legislation, the Prevention of Food Adulteration Act, seeks to test only end products, and does not ensure the adoption of the principles of HACCP throughout the food chain.

The new enactment – the Food Safety and Standards Act, 2006, though not operational – seeks to incorporate HACCP principles. In seeking to consolidate these legislative instruments into one combined whole, it is a serious attempt at harmonizing legislation to comply with international standards. Some flaws in the legislation may be pointed out here. Both the organized as well as the unorganized food sectors are required to follow the same food law. The stringent norms relating to specifications, traceability and recall procedures are also extended to the informal food economy in the country. This may adversely affect street food sellers and stalls. The act excludes plants prior to harvesting and animal feed from its purview and hence does not control the entry of pesticides and antibiotics into the food at its source.

2.3. Animal health

2.3.1. Legislation

Among the pieces of central legislation the following are the main ones:

*Wild Life (Protection) Act, 1972*

The Wild Life (Protection) Act seeks to protect wild animals, birds and plants with a view to ensuring ecological and environmental security. Although this enactment does not specifically deal with the issue of animal health, two specific sections dealing with the preventive aspects of wildlife
health are worth noting. Section 32 states that no person shall use chemicals, explosives or any other substances which may cause injury to or endanger any wildlife in any wildlife sanctuary. Section 33A, introduced by an amendment to the act in 2000, mandates that the Chief Wildlife Warden shall take measures for the immunization against communicable diseases of livestock kept in or within five kilometres of a sanctuary.

Livestock Importation Act, 1898

The Livestock Importation Act, which was amended in 2001 by the Livestock (Importation) Amendment Ordinance, provides for the regulation of the import of livestock which is liable to be affected by infectious or contagious disorders. The central government may regulate, restrict or prohibit any stock which may be liable to be affected by infectious or contagious disorders and any fodder, drug, stable-litter, clothing harness or fittings appertaining to livestock (sect. 3). The act empowers customs officials to act as though empowered under Section 11 of the Customs Act, 1962.

Section 3-A specifically states that the central government may by notification "regulate, restrict or prohibit in such manner and to such extent as it may think fit, the import into the territories to which this act extends or any livestock product, which may be liable to affect human or animal health."

The act empowers the state governments to make rules on the detention, inspection, disinfection or destruction of imported livestock and other items as well as on the powers and duties of those they appoint.

2.3.2. Institutions

The Ministry of Environment and Forests (MEF) and the Ministry of Agriculture (MOA) are the key ministries in charge of animal health concerns regarding domesticated animals. The Department of Animal Husbandry and Dairying has been given the task of monitoring and coordinating the various institutions that are engaged with animal health. MEF is entrusted with the task of protection of wildlife health in sanctuaries and wildlife parks. Each state government has the power to protect the health of animals within its own boundaries and has been empowered by state enactments to set up quarantine stations and to test for diseases. In case epidemic outbreaks, the central government issues
notifications and guidelines to control and monitor the disease, and has in several instances set up ad hoc monitoring committees.

The mandate of the animal quarantine and certification services within the MOA is to prevent the entry of livestock diseases into India by regulating the import of livestock and livestock-related products, and providing export certification for livestock and livestock products which are exported from India.

In order to provide referral services over and above the existing disease diagnostic laboratories in the states, one central and five regional disease diagnostic laboratories have been set up to strengthen the existing facilities. The Centre for Animal Disease Research and Diagnosis of the Indian Veterinary Research Institute, Izatnagar, is functioning as the central laboratory.

2.3.3. Evaluation

With regard to animal health, there is a need for a more effective centralized authority to monitor and coordinate the various activities of the state authorities. More effort at border control and monitoring is also needed. Further, there is need for a more sustained effort to ensure that the wildlife protection laws are strengthened to ensure protection of wildlife parks and sanctuaries and wildlife habitats.

2.4. Plant health

2.4.1. Plant quarantine legislation

_Destructive Insects and Pests Act, 1914_

The Destructive Insects and Pests Act is a pre-independence law which continues to regulate the introduction and movement of any insect, fungus or pest which could be destructive to crops. It has gone through several amendments over the years.\(^9\)

Under the act, the central government can prohibit or regulate the import into India of any insects or articles (or classes thereof) likely to cause

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\(^9\) See Destructive Insects and Pests (Amendment) Act, 1930; Destructive Insects and Pests (Amendment) Act, 1938; Destructive Insects and Pests (Amendment) Act, 1939; Destructive Insects and Pests (Amendment and Validation) Act, 1992.
infection to crops, by issuing a notification in the gazette (sect. 3(1)). The act further empowers the government to regulate the transport of insects or articles likely to cause infection to crops from one state in India to another (thus providing for domestic regulation) (sect. 4(a)). The act also empowers state governments to make rules for specific purposes in order to aid the central government in fulfilment of the main tasks of preventing the spread of these pests (sect. 4(a)).

*Plant Quarantine (Regulation of Import into India) Order, 2003*

With this new Plant Quarantine Order, agricultural imports into India are now classified into one of the following categories and have to follow these procedures for import:

- **prohibited plant species:** These are plants/planting materials and countries from which import is prohibited. Justifications for the same are listed in Schedule IV (cl. 3(2));

- **restricted species:** These are plants and plant materials the import of which into India is restricted and permissible only with the recommendation of an authorized institution and an import permit with an additional declaration and special conditions as provided under Schedule V of the order (cl. 10(1)). Phytosanitary certification has to accompany the consignment as well (cl. 10(2));

- **species requiring additional declarations and special conditions:** The same as above except that no recommendation is required from issuing authorities; and

- **plant material imported for consumption or industrial processing:** These are plants/planting materials for which imports are permissible on the basis of a phytosanitary certificate, an inspection conducted by inspection authority and treatment as may be required (cl. 3(1)).

As per clause 14(1) of the order, the central government, through the Joint-Secretary in charge of Plant Protection in the Department of Agriculture and Cooperation, can relax any of the conditions of this order in the public interest. The powers for relaxing conditions of import permits and phytosanitary certificates for one-time exception have been delegated to officers in charge of plant quarantine stations.
(a) Permits

The notable feature of the order is that it has brought about a strict permit regime. An import permit is rather simply defined as "an official document authorizing the importation of a consignment in accordance with specified phytosanitary measures" (cl. 2(x)). No consignment of items regulated under the order is allowed into the country without a valid permit (cl. 3(1)).

Valid import permits can only be issued by the permit-issuing authorities, which are listed in Schedule X of the order. Distinct import permits are to be issued for special products, e.g. live insects and microbial cultures (cl. 7) and germplasm, transgenic or GMOs (cl. 6).

(b) GMOs

The order also seeks to regulate the import of GMOs of plant origin for the purpose of agricultural research or experimentation (cl. 6(1)-(3)). Such an import would require a permit to be issued by the Director, National Bureau of Plant Genetic Resources (cl. 6(1)). These permits will be issued subject to the approval of the Genetic Engineering Approval Committee (GEAC) or the Review Committee on Genetic Manipulation (RCGM), as the case may be (cl. 6(2)). However, the order clearly provides that this does not cover imports for commercial purposes, which are governed by separate clearances.11

(c) IPPC: compliance and derogation

The order purports to promote harmony with the International Plant Protection Convention (IPPC)12 through the following:

- phytosanitary measures under the order are to be based on justified scientific principles with pest risk analysis (PRA) as their cornerstone. The definition adopted for PRA is the same as that in

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10 See Section 1.7.
11 Clause 8(3) provides that “bulk shipment(s) of transgenic plants or plant products or genetically modified organisms shall be dealt with per the provisions of the Rules for manufacture, use, import, export and storage of hazardous micro-organisms, genetically engineered organisms or cells made under [Sections 6, 8 and 25 of the Environment (Protection) Act]”.
12 See Chapter 2, Part III for an explanation of the IPPC.
the IPPC. As per clause 3(7), the guidelines for PRA have to be based on the standards established by the IPPC;
• the inspection and certification provisions (cl. 3, 8 and 10) under the order are in compliance with the requirements of article IV of the IPPC;
• under the definitions in the order, phytosanitary certificates are defined as "certificates issued in the model format prescribed under the IPPC and issued by an authorised officer at country of origin of consignment or re-export" (cl. 1(xix)). Article V of the IPPC is complied with in this regard;
• the restriction placed on the entry of certain plants and planting material by the order (cl. 3(14)) is in compliance with requirements for the same under the IPPC (art. VII (2)(d));
• the order is freely accessible to all, with a copy being available on the website of the national plant protection organization; and
• as per the notifications issued by the WTO Committee on Sanitary and Phytosanitary Measures, the order is "in line with the International Standards of Phytosanitary Measures of the [IPPC]".13

Plant Quarantine Bill, 2004

The Plant Quarantine Bill sought to establish the Plant Quarantine Authority of India (PQAI). The PQAI would be specifically established to meet India’s obligation under the IPPC to establish a central regulatory agency for plant protection, a national plant protection organization. The bill seeks to bring about a comprehensive regulatory framework for prevention of the spread of quarantine pests both domestically as well as outside national boundaries. The bill seeks to finally repeal the Destructive Insects and Plants Act.

2.4.2. Pesticide legislation

Insecticides Act, 1968

Another relevant piece of legislation regarding plant health is the Insecticides Act and the rules framed thereunder. This legislation and its rules seek to ensure the availability of quality, safe and efficacious pesticides to the farming community and to manage risks to human health and the environment.

13 WTO Committee on Sanitary and Phytosanitary Measures Notification No. G/SPS/N/INDIA/12, 4 March 2004.
The act seeks to regulate the import, manufacture, sale, distribution, use and transport of insecticides (including herbicides, fungicides, rodenticides, etc.). The Ministry of the Agriculture (MOA) is the relevant ministry under the act. The Central Insecticides Board and Registration Committee along with the Directorate of Plant Protection, Quarantine and Storage in the MOA are the authorities concerned with the registration requirements and other related matters.

2.4.3. Seed legislation

*Seeds Act, 1966*

The relevant Indian enactment for seeds is the Seeds Act. This act provides for the regulation of the quality of only certain seeds, which are to be notified by the central government (sects. 5, 7). The main institution brought into being by this act is the Central Seeds Committee, which is constituted by the central government (sect. 3(1)). The primary function of this committee is to advise the central and state governments on matters arising out of the administration of this act (*id.*).

A relevant aspect to be kept in mind with regard to this act is that authorities created under it are entitled to act only in the case of seeds sold for agricultural purposes and not for human consumption. The relevant enactment for the latter is the Essential Commodities Act, 1955.

*Seeds Bill, 2004*

The Seeds Bill, 2004, is proposed as a replacement for the Seeds Act, 1966. As per Section 12 of the bill, all kinds and varieties of seeds have to be registered in the National Register of Seeds. No seed can be sold (for the purpose of planting) unless it is registered (sect. 13). The designated body for registration is the registration sub-committee (which comes under the Central Seeds Committee) (sect. 12).

One of the most controversial and for our purposes relevant provisions of the Seeds Bill is Section 15 which provides in effect for registration of transgenic seeds under the bill and as a result thwarts existing biosafety regulations.
For Biosecurity purposes, Section 18 provides the grounds for exclusion of certain varieties of seeds from registration. The grounds for such exclusion are if:

- "prevention of commercial exploitation of such kind or variety is necessary to protect public order or public morality or human, animal or plant life and health, or to avoid serious prejudice to the environment" (sect. 18(1)); and
- it is "a kind of variety of seed containing any technology, which is harmful, or potentially harmful" (sect. 18(2)).

Section 36 of the bill deals with the import of seeds and it provides for the compulsory registration of all imported seeds (although the government may allow the import of an unregistered seed for research purposes). Further, all imports of seeds "shall be subject to the provisions of the Plants, Fruits and Seeds (Regulating of Import into India) Order, 1989, or any corresponding order made under Section 3 of the Destructive Insects and Pests Act, 1914".

2.4.4. Evaluation

Some basic themes emerge in an analysis of the plant quarantine framework in India. The Destructive Pests and Insects Act, 1914, along with the Plant Quarantine Order, 2003, seek to deal with this rather complicated issue. In certain areas there are obvious shortcomings while in others the current set-up can be said to be a success.

The obvious shortcomings of the Destructive Pests and Insects Act, 1914, are that its definition of plant protection is limited to crops – defined to include all agricultural and horticultural crops and all trees, bushes or plants – which leaves out any sort of protection for other areas, e.g. forests.

None of the enactments deal with the issue of exports and phytosanitary certification for exports. Thus, in case of exports the requirement of phytosanitary certification is not mandatory. This has resulted in cases where exporters have ended up exporting articles without seeking the requisite certification, due to an unawareness on their part of such a facility existing or an unwillingness to obtain the same. Some consignments have been returned, causing a loss of faith in Indian exports. Under the current set-up, officers notified under Notification 8-97/91-PP.I issued by the Ministry of Agriculture
(Department of Agriculture and Cooperation) on 26 November 1993, are authorized to inspect, fumigate or disinfect and grant a phytosanitary certificate.

The fact that the existing certification process might not be performing adequately is clear from the circular issued by the Ministry of Agriculture to the certificate-issuing authorities in May 2006, which pointed out a number of cases where although phytosanitary certifications had been issued by such authorities to certain consignments, these consignments had been rejected by the countries of import on phytosanitary grounds. This theme of non-compliance with the existing framework and inability of the existing machinery to follow the letter of the law runs throughout India’s Biosecurity-related legislation and the regulatory framework it creates.

With regard to monitoring imports of regulated articles, the frequent updating of the Plant Quarantine Order, 2003, suggests that the concerned department prioritizes this regulatory area. However, India does not seem to have put in place an adequate mechanism. For the system to work with a certain degree of competence, it has to put in place a paperless system that feeds into the existing national network of connected computer servers for customs purposes. A comprehensive border monitoring mechanism should also be put in place.

2.5. Invasive alien species

2.5.1. Legislation

The enactment of the Biological Diversity Act, 2002, was necessitated by virtue of India’s signing and ratifying the Convention on Biological Diversity (CBD). Though the CBD provides sufficient latitude to its members to pursue distinct approaches to national biodiversity laws, India chose to adopt the route of having stand-alone legislation on biodiversity.

With regard to Biosecurity, the Biological Diversity Act, 2002, only has limited relevance. To begin with, there is no provision in the act to deal with invasive alien species (IAS). In fact, no mention is made of these species throughout the legislation.

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15 See Chapter 2, Part VI for a full description of the CBD.
With regard to living modified organisms (LMOs), Chapter IX contains a very general provision which encumbers the central government to take measures "to regulate, manage and control the risks associated with the use and release of living modified organisms resulting from biotechnology likely to have adverse impact on the conservation and sustainable use of biological diversity and human health" (sect. 36(4)(ii)).

Apart from these provisions, rather general duties are imposed upon the central government to develop strategies, plans and programmes for the "conservation and promotion and sustainable use of biological diversity" (sect. 36(1)) and to integrate these goals of conservation and sustainable use of biological diversity into "relevant sectoral, and cross-sectoral plans, programmes and policies" (36(3)).

Under Section 38, the central government may also notify certain threatened species and "prohibit or collection thereof for any purpose and take appropriate steps to rehabilitate and preserve those species". Finally, Section 40 gives the central government the power to exempt certain biological resources from the provisions of the act, including "biological resources normally traded as commodities".

2.5.2. Institutions

The Biological Diversity Act, 2002, sets up a whole institutional framework for the protection and sustainable utilization of biodiversity in the country. These include the National Biodiversity Authority, State Biodiversity Boards in every state and Biodiversity Management Committees at local levels. This three-tier institutional framework and the relevant roles and responsibilities are further dealt with and elaborated in the Biological Diversity Rules, 2004.

2.5.3. Evaluation

The lack of adequate domestic regulation to protect biodiversity is an issue of great concern. The seriousness of the problem is compounded by the fact that India is a biodiversity-rich country with numerous agro-economic zones. The lack of domestic regulation is often blamed on the unwillingness of the state governments to comply with any strict regulations in this regard and the inadequacy of the existing enforcement machinery.
The issue of IAS for forest areas is not dealt with under the regulatory framework in place. The general view seems to be that this issue is a concern of the Ministry of Environment and Forests (MOEF) and should be dealt with by that ministry (possibly under the set-up created by the Biological Diversity Act).

2.6. Biosafety

For biosafety, the regulatory framework consists of rules issued in 1989 by the MOEF under the Environment Protection Act, 1986. These have been revised by guidelines issued in 1990, 1994 and 1998 (issued vide Rule 4(2) of the aforementioned rules). The fact that these were brought in place even before the Rio Summit in 1992 which adopted the CBD shows that India was one of the pioneers in this regard.

The 1990 Recombinant DNA Safety Guidelines and the 1994 Revised Guidelines for Safety in Biotechnology contain detailed guidance on containment and safe laboratory practices for GMOs in both the agricultural and pharmaceutical sectors. The 1998 Revised Guidelines for Research in Transgenic Seeds, Plants and Plant Parts, on the other hand, apply only to GMOs used in the agricultural sector.

The 1990 guidelines made one fundamental change from the 1989 rules vis-à-vis their treatment of the deliberate treatment of GMOs. Whereas such a release was permitted only under special circumstances under the rules (para. 9(1)), the guidelines permit them while focusing on assessing and managing possible environmental and health risks (para. 9).

2.6.1. Institutions

These rules and guidelines have put in place "multi-layered decision-making structures". What this means in practice is a multitude of bodies which come under two different ministries. The structure involves six different bodies which come into play over the four different phases a biotechnology product or organism has to undergo.

The first phase is pre-research, where the appropriate body is the Recombinant DNA Advisory Committee, which is constituted by and based in the Department of Biotechnology (DBT) of the Ministry of Science and Technology and is in charge of giving pre-research approvals. The second phase is the research phase for which the appropriate authority is the RCGM,
which is also constituted by and based in the DBT and which is charged with monitoring the research and experimental release of biotechnology products and organisms. A monitoring and evaluation committee (MEC) comprising scientists, agricultural experts and other officials nominated by relevant ministries has been formed under the RCGM.

The next phase is commercial release, which comes under the purview of the GEAC, which is constituted by and based in the MOEF and gives approval for such release from an environmental perspective. The last phase is post-release which involves the MEC, the State Biotechnology Coordination Committee and the District Level Committee. Apart from this, the Institutional Biosafety Committee is charged with implementing and monitoring safeguards at the research and development sites (under the supervision of the post-release-phase bodies).

2.6.2 Legislation

Shift from case-by-case to event-based approval

Until June 2006, the GEAC was following a "case-by-case" approval process for genetically modified (GM) crops. Under this system, every GM hybrid/variety had to undergo a minimum of three years of official trials before being approved. On 30 June 2006 as per a decision of the GEAC, an "event-based approval system" has been put into place instead, which is supposed to speed up the whole process. An "event" refers to a specific gene construct that can be incorporated in a number of existing hybrids or varieties.

Import of GM products

On 7 April 2006, the regulation of importation of GM products was provided for under the Foreign Trade Policy, 2004–2009. MOCI, through the Directorate General of Foreign Trade, notified new regulations for import of GM products by amending Schedule I (Imports) of the ITC (HS) Classification of Export and Import Items under Section 5 of the Foreign Trade (Development and Regulation) Act, 1992. As a result of this notification:

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16 Decision taken in the 69th meeting of the GEAC held on 30 June 2006, available at www.envfor.nic.in
The import of GMOs/LMOs for the purpose of (i) R&D; (ii) food; (iii) feed; (iv) processing in bulk; and (v) for environmental release will be governed by the provisions of the Environment Protection Act, 1986, and Rules, 1989.

The import of any food, feed, raw or processed, or any ingredient of food, food additives or any food products that contain GM material and are being used either for industrial production, environmental release or field application will be allowed only with the approval of the GEAC.

Institutes/companies who wish to import GM material for R&D purposes will submit their proposal to the RCGM under the DBT.17

Crucially, it is further provided that all GM consignments have to carry a declaration to that effect at the time of import, with provision for penal action under the Foreign Trade (Development and Regulation) Act, 1992, in case of non-compliance.18 These conditions were, however, kept in abeyance for three months via a notification issued by the Director General of Foreign Trade on 4 May 2006.19 The United States filed notifications with the WTO the same month against this regulation,20 seeking clarifications about the amendments and hinting at initiating action against India under the TBT and SPS Agreements.

Apart from this, the provisions of the Plant Quarantine Order, 2003, are applicable for the import of transgenic seeds (not for commercial purposes).21

17 Condition 18(a), (b) and (c) of Chapter 1A (General Notes Regarding Import Policy), Schedule-I (Imports) of the ITC (HS) Classifications of Export and Import Items, 2004–2009, inserted vide Notification No. 2 (RE-2006)/2004-2009, New Delhi, 7 April 2006, available at exim.indiamart.com.
18 Id. Condition 18(d). This offers the crucial distinction between the 1989 Rules and these conditions, since such a declaration at the point of entry was totally voluntary under the rules, see Decision taken in the 66th meeting of the GEAC held on 2 May 2006, available at www.envfor.nic.in.
20 G/TBT/N/IND/12, 17 May 2006 and G/TBT/N/IND/17, 23 May 2006.
21 See Section 1.5.1.
2.6.3. Evaluation

Though the existing rules and guidelines seek to delineate the various functions of the institutions in place, certain grey areas exist. Thus, while RGMC is supposed to administer experimental research and the GEAC supervises the deliberate release of transgenic crops, the question arises regarding under which function field trials would fall. Public interest litigation filed by a non-governmental organization forced amendments to the 1998 Biosafety Guidelines in September 1999 to the effect that the RCGM is now authorized to approve small experimental field trials for research.

A serious shortcoming of the existing regulatory set-up is that it fails to take into account other existing legislation concerning biotechnology. This includes: (a) the Seeds Act; (b) the Biosecurity Regulations; (c) the Biodiversity Act; (d) the Protection of Plant Varieties and Farmers Rights Act; and (e) the Prevention of Food Adulteration Act.

To replace the GEAC with an autonomous statutory body, a National Biotechnology Regulatory Authority, along the lines of India’s Atomic Energy Regulatory Board, is under discussion. The recommendation to create this authority was first made by the Task Force on Agricultural Biotechnology (chaired by M.S. Swaminathan) in its report of May 2004.22 This call was repeated in the National Biotechnology Development Strategy, which was prepared by the DBT in 2005.23 However, it must be borne in mind that no such demand for reform had emanated from the MOEF, which is the ministry responsible for the GEAC.

Some critical aspects need to be kept in mind while evaluating India’s legal regulatory setup for biosafety vis-à-vis the requirements under the Cartagena Protocol on Biosafety.24 India’s existing regulatory framework is considered to be strict and one that provides for all adequate safeguards. This has meant that India has not been required to reform this set-up in order to bring about compliance with the Cartagena Protocol. The coming into force of the protocol has been considered an event that legitimizes the existence of the present framework. However, it must be pointed out that current Indian law does not provide any procedure for an advance informed agreement.

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22 This task force was set up by the Ministry of Agriculture. See Task Force Report, supra note 1.
23 See National Biotechnology Development Strategy, Department of Biotechnology - Ministry of Science and Technology, Government of India, launched on 31 March 2005.
24 For a discussion of this instrument, see Chapter 2, Part VII.
The stringent nature of the regulatory framework when compared with international standards can be gauged by the requirement of agronomic analysis (socio-economic analysis) to be a part of the procedure of risk assessment (along with the usual ecological and human health safety evaluations). This requirement is unique and is in addition to any framework generated solely under the Cartagena Protocol.

There is broad agreement that the aspect of biosafety that requires close inspection and lengthy deliberation concerns the ability to actually bring into effect the regulatory mechanism put in place on paper. There are three shortcomings in the Indian context in this regard: (i) the basic lack of technically trained manpower and adequate machinery (both quantitatively as well as qualitatively); (ii) lack of interest in strictly enforcing the laws in place. The regulatory framework tends to prefer being pragmatic (in the sense of flexibility) rather than being strict, a tendency that can be noted in other areas examined in this chapter as well. It appears that extraneous concerns weigh heavily on decisions as to enforcement of the regulatory system. (iii) There is also a perceptible lack of coordination in the system in place, with various ministries contending for a greater role in the process.

Of particular relevance for the previous point is that the Biosafety Clearing-House mechanism provided for under article 20 of the Cartagena Protocol has been established and is functioning in India. In this regard the MOEF is currently implementing a Global Environmental Facility/World Bank-funded project on capacity building in the context of the protocol. One of the areas where capacity is sought to be developed in this context is the strengthening of the legislative framework and operational mechanisms.

### III. CONCLUSIONS

This analysis of the Biosecurity legal framework has been undertaken applying the FAO definition of Biosecurity. The primary elements that constitute Biosecurity cover the introduction of plant pests, animal pests and diseases and zoonoses, the introduction and release of GMOs and their products and the introduction and management of IAS and genotypes.

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The concept of *Biosecurity* being nascent, evolving as it is with progress in science and technology, it has not been incorporated as an integrated whole into legislation in India. So the approach here is essentially piecing together sectoral pieces of legislation that have a different historical background, in an attempt to test their feasibility against emerging concerns around *Biosecurity*. At the outset, therefore, it is important to acknowledge this limitation and the essential pitfalls in rereading the enactments with a different prism.

The *Biosecurity* legal framework of India is presently evolving. The existing framework on sectoral issues relating to *Biosecurity*, both on the statute books and the institutional structures, is both disparate and elaborate. This review sets out to map this elaborate framework, keeping in mind the historical context and continuing relevance. It also alludes to the proposed changes to the existing framework and the newer pieces of legislation that are on the drawing board of the relevant legal departments.

The challenges for implementation of the *Biosecurity* regime in India are immense, given the size and geographical variations within the country. Lack of trained manpower and the resources for scientific research are additional challenges that loom large. In some of the other countries that have undertaken a similar exercise, there is a suggestion to consolidate existing legislation and create a single agency to deal with *Biosecurity* concerns. However, this approach needs more careful consideration in the Indian context. The motivations behind the existing legal framework and the focus of work of the respective institutions differ vastly. Besides, the *Biosecurity* concerns do not necessarily override the pre-existing purposes behind the sectoral legislative instruments and the institutions set up under them. An altogether new legal framework, with institutions tailored to carry out the tasks of protecting and promoting *Biosecurity* within the delimitations of their respective mandates, could perhaps be a more effective approach.

It may be stated that currently, there is no clear indigenous understanding of the concept of *Biosecurity*. The draft National Policy for Farmers, put together by the National Commission on Farmers, refers to a "National Agricultural *Biosecurity* System", which discusses the concept at some length. The approach contained in this document is narrower than the definition adopted by FAO in its COAG document.27

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27 See id.
More importantly, the concept of Biosecurity needs to be viewed more broadly from the perspective of public policy on health, environment and sustainable development. Evolving international standards are driven by interests that may not be consistent with a broader Biosecurity approach.

The various standards that are being prescribed to ensure Biosecurity provide a broad template for compliance. However, the politics behind the standard setting are of equal importance. Standards and technical regulations for Biosecurity may be viewed from the two different intents with which they are put in place. The two primary purposes are: the promotion of trade, and the promotion of public policy objectives. Although there are several fundamental differences between them, they both depend on the same quality assurance institutions and are governed by many of the same legal regimes. Although many of the weaknesses that exist in these institutions and legal regimes do not create problems in the context of trade promotion, they do create problems in the context of public policy promotion.

Finally, it is important that the focus of legislation, including legislation dealing with Biosecurity concerns, be directed towards protecting and conserving the environment, and ensuring the health of the country’s people, flora and fauna. While trade concerns are important and should run a parallel course, there is an urgent need to keep the focus on the broader concerns as expressed in CBD and the Rio Declaration, particularly the fundamental rights to clean environment, food, health and life.
KENYA COUNTRY STUDY*

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* This chapter was prepared by Patricia Kameri-Mbote.
I. INTRODUCTION

This chapter is divided into three parts. After this introduction, part II lays out the normative and institutional framework for Biosecurity in Kenya. It looks at the constitutional basis for Biosecurity, SPS-related and food safety laws and institutions and laws on biodiversity and biosafety.

Part III analyses the adequacy of the Biosecurity framework and notes that there are overlaps that need to be addressed. It is clear that Kenya has considered the need to synergize the Biosecurity framework. The establishment in 1996 of the Kenya Plant Health Inspectorate Service (KEPHIS), a parastatal agency under the Ministry of Agriculture, began a process of bringing the phytosanitary aspects of Biosecurity under one rubric. This process is still ongoing. Similarly, the promulgation of a framework environmental law, the Environmental Management and Coordination Act (EMCA) in 1999, and the establishment of the National Environment Management Authority (NEMA) as a coordinating institution under it, is a further step towards coordinated performance of Biosecurity functions.

It is, however, a matter of concern that the Biosafety Bill is still in draft form and genetic modification activities in Kenya have been proceeding only on the basis of draft regulations prepared in 1998. Part IV comprises a conclusion and recommendations where the basic issues for consideration in reviewing and framing a national Biosecurity legal and institutional framework in Kenya are outlined.

II. BIOSECURITY LAWS IN KENYA

Although Kenya has a host of statutes dealing with Biosecurity, it is notable that the purposes of these acts are as varied as the acts themselves. The framework environmental law, the EMCA, for instance, establishes the legal and institutional framework for the management of the environment and for matters connected and incidental thereto as its main purpose. The Biosafety Bill for its part has as its main objective ensuring an adequate level of protection in the field of safe transfer, handling and use of genetically modified organisms (GMOs) resulting from modern biotechnology that may have an adverse effect on the environment; and to establish a transparent and predictable process to review and make decisions on such GMOs and related activities. While it is not clear whether the GMO bill is promulgated under
EMCA, it is clear that EMCA’s mandate extends to biosafety. In discussing the legal framework, therefore, we divide the laws into three categories.

The first category comprises laws that provide the context within which Biosecurity occurs. This includes the Constitution. The second category addresses SPS-related and food safety laws. The third category analyses laws related to biodiversity management and biosafety.

2.1. Constitutional anchorage for Biosecurity

Kenya’s Constitution does not contain explicit Biosecurity provisions. It does, however, place importance on the right to life, and experts argue that the right to life encompasses the right to a clean and healthy environment.\(^1\) It protects individual fundamental rights and freedoms which are relevant to ensuring the integrity of biological resources and food safety. These include the right to life and the right to the protection of the law, which appear in Chapter V of the Constitution. These rights provide the necessary context for implementing Biosecurity. It is instructive to note that the Constitution also provides for the right to sue and therefore provides a means of ensuring Biosecurity through legal interventions.

The stalled constitutional review process was expected to define a more explicit constitutional basis for securing environmental integrity. All the drafts generated during the process explicitly required the state to "ensure the respect and integrity of natural processes and ecological communities, including conservation of habitats and species; ensure sustainable exploitation, utilization, management and conservation of the environment and natural resources and the equitable sharing of the accruing benefits; and prevent pollution and ecological degradation" (draft Proposed National Constitution of 2005, sect. 87). The drafts also included explicit provisions on the right to a healthy environment. Section 88 provides that every person has a duty to:

- (a) ensure ecologically sustainable development and use of natural resources;
- (b) respect, protect and safeguard the environment;
- (c) prevent or discontinue an act which is harmful to the environment;

(d) direct the appropriate authority to take measures to prevent or
discontinue an act or omission which is harmful to the environment;
and
(e) maintain a clean, healthy and safe environment.

Section 89 on conservation of the environment provides that in the
utilization and management of the environment, the state shall protect
genetic resources and biological diversity; establish systems of environment
impact assessment, environment audit and monitoring of the environment;
and ensure that the environmental standards enforced in Kenya are in
harmony with accepted international standards. An institutional framework,
the National Environment Commission, was also provided for in the drafts
to oversee the implementation of these provisions (sect. 92). All these
provisions have implications for *Biosecurity* in Kenya and provide the context
within which it is anchored.

Even though the constitutional review process is yet to be completed, it is
noteworthy that the provisions on the environment are not among the
contested ones. It is expected that when the review process is completed,
these provisions will be part of a new Constitution.

### 2.2. Sanitary- and phytosanitary-related legislation

Sanitary and phytosanitary (SPS) measures are measures that protect human,
animal and plant health. Kenya has a number of laws that deal with these
measures by regulating animal and plant health, food safety and related issues
of packaging, importation, exportation, manufacture, distribution and use
thereof. The following laws are in place:

- Seeds and Plant Varieties Act (Chapter 326), 1972;
- Plant Protection Act, (Chapter 324), 1962;
- Pest Control Products Act, (Chapter 345), 1983;
- Suppression of Noxious Weeds Act, (Chapter 325), 1986;
- Animal Diseases Act, (Chapter 364), 1972; and
- Cattle Cleansing Act, (Chapter 359), 1937.

One discernible factor in these laws is the overlapping mandates of
regulatory agencies. Indeed, in 2003, an attempt was made to come up with

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one law amalgamating the Seeds and Plant Varieties Act, Plant Protection Act and the Suppression of Noxious Weeds Act. This idea was abandoned but it is expected that once the revision of the Seeds and Plant Varieties Act is completed, the other statutes will need to be amended to align their objectives with those of this act as well as to consolidate the roles of KEPHIS in the phytosanitary area.

2.2.1. Plant health

*Seeds and Plant Varieties Act (Chapter 326), 1972*

The Seeds and Plant Varieties Act deals with, among other things, the imposition of restrictions on the introduction of new seeds and plant varieties; the importation of seeds; and the prevention of injurious cross-pollination.

The regulatory agents under the act include the minister for the time being responsible for agriculture, seed analysts and KEPHIS. Established under the provisions of State Corporations Act (Chapter 446), 1996, KEPHIS is responsible for implementing phytosanitary and quarantine measures. It is the national focal point for SPS. The roles of KEPHIS include:

- coordinating all matters relating to crop pests and disease control;
- establishing service laboratories to monitor the quality and levels of toxic residues in plants as well as their soils and produce;
- advising the Director of Agriculture on appropriate seeds and planting materials for import and export;
- undertaking inspection, testing, certification, quarantine control, variety testing and description of seeds and planting materials;
- undertaking grading and inspection of plants and plant produce at the ports of entry and exit;
- enforcing standards for good husbandry and the control of pests and diseases;
- developing and implementing standards on both imported and locally produced seeds;
- approving all importation and exportation licences for plants and seeds issued by the ministry responsible for commerce and industry before such importation or exportation is implemented;
• implementing the national policy on the introduction and use of genetically modified (GM) plant species, insects and micro-organisms and regulating imports of GM seeds; and
• establishing posts at convenient locations for quarantine, inspection and quality control of fertilizer and seed, and monitoring agricultural inputs and their environmental effects.

To enable it to discharge its duties under the bill, KEPHIS is granted the power to appoint seed inspectors, seed analysts and plant examiners.

With respect to seeds, the minister is required to consult representatives of the organizations having a substantial interest in the matter being regulated (sect. 3). Seed regulations under both the draft bill and the current act seek to control and regulate seed production, processing, testing, certification, and marketing by:

(a) ensuring that reliable and adequate information is afforded as to the nature, condition and quality of seeds intended for sale;
(b) preventing the sale of seeds which are deleterious, which have not been produced in specified conditions, which have not been tested for purity or germination or which are of a variety of which the performance has not been subjected to trials;
(c) requiring the registration and/or deregistration of persons growing any specified crop for the main purpose of seed production, or of persons selling any seed;
(d) supporting plant inspectors in preventing the spread of plant diseases by seeds;
(e) providing for seed certification, processing, sampling, testing and marketing;
(f) regulating the descriptions under which seed is sold;
(g) regulating, controlling or prohibiting the import or export of seeds;
(h) charging fees for services rendered by KEPHIS under the act or the Seed Regulations;
(i) prescribing the national obligations for seed crops; and
(j) providing for the filing of appeals.

These regulations may also address packaging, bags, trays or other containers in which seeds may be sold as well as the requirements for the marking of such containers; prohibit the selling of uncertified seeds; and require seed dealers to maintain records on seed transactions. Furthermore, KEPHIS has power
under the draft bill to limit the importation of seeds in certain situations, such as where the seeds are unsuitable in Kenya because they are of a type developed in countries with different climatic or other conditions (sect. 15).

In order to maintain the purity of seeds, KEPHIS is further given the authority under the draft bill to restrict the growing of seeds in any area in Kenya in order to isolate them from crops that might cause injurious cross-pollination.

**Plant Protection Act (Chapter 324), 1962**

The Plant Protection Act makes provision for the prevention of the introduction and spread of diseases destructive to plants. Pests are defined as any animal or vegetable organism inimical to the growth or existence of living plants or injurious to plant products and any other agent capable of producing a communicable disease of plants (sect. 2). The main regulatory agents under the act are the minister for the time being responsible for agriculture and inspectors who comprise the Director of Agriculture and any other persons authorized by the director, mainly from KEPHIS.

Under the act, inspectors are authorized to enter any land or building other than a dwelling house for the purpose of discovering pests or diseases in any plants, upon informing the occupier or owner (sect. 5).

The minister can order a prohibition or restriction or can regulate the importation and exportation of any plants and the soil, packages, coverings or wrappings thereof and any other article, animal or insect likely to infect any plant with any pest or disease. In carrying out this role, the minister is authorized to and has made rules for a variety of measures which can be taken (sect. 3).

Owners and occupiers of land are obligated to take the measures required by the minister and which are reasonably necessary for eradicating, reducing or preventing the spread of any pest or disease (sect. 4). Persons who knowingly introduce any pest or disease into any cultivated land or who wilfully obstruct or interfere with an inspector are guilty of an offence (sect. 7).

The minister may, if he or she thinks fit, order compensation to be paid out of public funds to any person whose plants or other articles are destroyed pursuant to the act (sect. 6). The main compliance mechanism established
under the act is through the use of criminal sanctions (sect. 9). Inspectors are insulated against any suit, prosecution or other legal proceeding for anything done in good faith and without negligence (sect. 10).

Pest Control Products Act (Chapter 345), 1983

The Pest Control Products Act regulates the importation, exportation, manufacture, distribution and use of products intended for the control of pests and of the organic function of plants and animals. Under Section 5, the act establishes the Pest Control Products Board (PCPB) whose functions include:

(a) assessing and evaluating pest control products;
(b) considering applications for registration of pest control products; and
(c) advising the minister on all matters relating to the enforcement of the provisions of the act and its regulations (sect. 6).

The enforcement of regulatory mechanisms established under the act is the responsibility of inspectors and analysts. Under Section 8 of the act, the minister responsible for agriculture is obligated to appoint suitably qualified persons as inspectors and analysts. Inspectors have the power to enter any place or premises for the purposes of carrying into effect any of the provisions of the act; in instances where they reasonably believe a pest control product to which the act applies is or has been manufactured, stored, sold or used; and where they reasonably believe there is material that is contaminated by a pest control product or which is used or capable of being used in the manufacture of a pest control product.

They also have power to examine any pest control product or material found in any place or premises or to open any package found in the premises which they believe contains any pest control product or material and take samples thereof; require any person to produce books, shipping bills, bills of lading, documents containing instructions or other documents or papers relevant to the performance of their duties, for the purpose of obtaining copies or extracts (sect. 9). Inspectors have the power to seize and detain any pest control product in the performance of their duties and may, under the orders of the court or with the consent of the person in possession of the products, dispose of the product (sect. 10).
Any person who refuses entry to an inspector acting under this section or obstructs him or her in making an entry or inspection or who, without reasonable excuse, fails to produce any pest control product or material for examination or any required document, is guilty of an offence (sect. 9(4)). The main compliance mechanism established under the act is criminal sanctions (sect. 12).

_Suppression of Noxious Weeds Act (Chapter 325), 1986_

The Suppression of Noxious Weeds Act provides that the minister of agriculture may by notice in the gazette declare a plant to be a noxious weed in any area, which may consist either of the whole of Kenya or of one or more districts or portions thereof.

Local authorities are empowered under Section 10 to make by-laws for securing the eradication of any noxious weed from land within their areas; for compelling owners or occupiers of land to cause any such weed to be eradicated from their land; and for such purposes, by-laws may appoint inspectors.

The act also creates the office of the inspector (a person authorized in writing by the Director of Agriculture to perform the duties of an inspector under this act in an area specified by the director, and a person appointed by or under by-laws made by the relevant local authority). The inspector may at all reasonable times enter upon land for the purpose of ascertaining whether any noxious weed or other weed exists thereon.

If an inspector finds upon land within a declared area any plant which has been declared to be a noxious weed, he or she may, by notice in writing to the person responsible for the land, require that person to clear the land of the noxious weed within a time specified in the notice. The notice shall state the particular noxious weed which has been found upon the land, and, as far as practicable, the portion or portions of the land on which the noxious weed has been found. If the person responsible fails to clear the land within the time specified in the notice, an inspector may, upon receiving a written authority from the director, enter, with or without assistance, upon the land and eradicate any noxious weed found there. Any expenses incurred in eradication shall be a civil debt recoverable summarily from the person responsible at the suit of the director. Failure to comply with this notice is an offence under the act.
2.2.2. Animal health

**Animal Diseases Act (Chapter 364), 1972**

The Animal Diseases Act provides for matters relating to the diseases of animals. It covers stock, ruminating animals, dogs, cats, rabbits and captive wild animals. Persons possessing animals infected with a notifiable disease (defined in Section 2 of the act) are required to keep the animal tied up in an enclosed place separate from uninfected animals.

The Director of the Veterinary Services Department (VSD) under the Ministry of Agriculture is to appoint inspectors for the purpose of implementing the provisions of the act. The director is empowered at Section 5 to declare any area "an area infected by a notifiable disease"; "to extend, diminish or alter ... an area declared to be an infected area"; to "declare an infected area to be free from a notifiable disease"; and to "prohibit the movement of animals from one ... area to another".

Once an area is declared to be infected:

- no stock shall be moved from or into it without the director's written permission;
- no animal shall be moved from the area unless it has been disinfected and treated in the manner required by the director;
- all stock in the area shall be herded as far as possible from any public road and shall not graze on any road reserve;
- the director may require the owner or person in charge of the animal to isolate the animal from other animals within the infected area or to remove the animal from the area;
- no person shall leave any such area without having complied with such reasonable precautions for preventing the spread of the notifiable disease as may be required by the veterinary officer or inspector in charge of the area; and
- the carcasses of all animals infected with the notifiable disease shall be disposed of in accordance with general or specific instructions issued by a veterinary officer or inspector.

Section 8 allows the director to prohibit for such time as he or she thinks necessary, or to regulate, the importation or exportation of all animals or any specified kinds of animals, or of carcasses, meat, hides, skins, hair, wool,
litter, dung, semen, live viruses capable of setting up infections in animals, sera, vaccines and other biological or chemical products intended to be used for the control of animal diseases.

Under Section 14, the director or inspector has power to enter any land, building, shed, van, truck or other premises or container to search for infected animals or their products such as meat, carcasses and hides. Subsidiary legislation under the act elaborates rules on issuance of permits, tests required and certification for importation and movement of animals; infected areas; and prevention of spread of diseases. Under the act, rules dealing specifically with birds; foot and mouth disease; rinderpest; and pig diseases have been promulgated. It is an offence punishable by law for any person to contravene the provisions of the act.

*Cattle Cleansing Act (Chapter 359), 1937*

The Cattle Cleansing Act provides for the cleansing of cattle. The responsibility for this is bestowed on the Minister of Agriculture as well as the Provincial Agricultural Board. Where the board has recommended to the minister that any area within that province should be a cattle cleansing area, the minister may, if he or she is satisfied that it will be of general benefit to the stock owners in the area, declare that area, or any part thereof, to be a cattle cleansing area.

Inspectors appointed under the act have the power and authority under Section 7 to inspect and count any cattle at any time. Further, a land owner in a cattle cleansing area on whose land cattle are kept is required to satisfy the inspector that there is adequate provision of facilities for the dipping or spraying of cattle and sufficient quantities of effective tick-destroying agents. It is unlawful under the act for any cattle owners to refuse or fail to submit their cattle for cleaning in accordance with the provisions of the act.

Under Section 10 of the act, where a land owner on whose land cattle are kept is absent from Kenya and does not have a representative or agent in Kenya with authority to carry out the terms of the act, any veterinary officer may authorize the due performance of these requirements. Any expenditure incurred in such circumstances shall be recoverable by the director as a civil debt.
2.2.3. Food safety

With respect to food safety, the relevant laws include:

- Food, Drugs and Chemical Substances Act (Chapter 254), 1970;
- Meat Control Act (Chapter 356), 1973;
- Standards Act (Chapter 496), 1974; and
- Public Health Act (Chapter 242), 1921.

Food, Drugs and Chemical Substances Act (Chapter 254), 1970

The Food, Drugs and Chemical Substances Act makes provision for the prevention of adulteration of food, drugs and chemical substances. Section 3 prohibits the sale of food containing anything poisonous or harmful; food that is unwholesome or not fit for human consumption, poisonous or which consists of filthy, putrid, disgusting, rotten, decomposed or diseased substances or foreign matter or which is adulterated. It is an offence under the act to label, package, treat, process, sell or advertise any food in a manner that is misleading or deceptive as regards its nature, value, substance, quality or composition (sect. 4).

Further, foods for which there are prescribed standards must conform to such standards, and labelling, packaging, selling or advertising such foods in a manner that misleads as to the conformity with the set standards is an offence (sect. 5). It is also an offence to sell, prepare, package, convey, store or display for sale any food under unsanitary conditions (sect. 7). The act makes it an offence to use or dispose of chemical substances in a manner likely to cause contamination of food or water for human consumption (sect. 24).

Subsidiary legislation under the act makes provisions for food hygiene, providing for licensing of premises used for sale, preparation, packaging, storage and display for sale of any food; prescribing conditions for growing and harvesting food; covering construction of food plants and facilities and other health measures to be taken in food plants. Subsidiary legislation under the act has addressed the issue of food labelling, additives and standards.

Meat Control Act (Chapter 356), 1973

The Meat Control Act provides standards for storage and transportation of meat and meat products intended for human consumption, slaughterhouses
and places where meat is processed, as well as import and export control over meat and meat products. Regulations made under the act require the licensing, control and regulation of slaughterhouses and premises where meat is processed for human consumption. They also specify standards to be observed, additives to be used and methods of packaging and labelling; require inspection of slaughterhouses and meat and meat products; and establish standards to be observed in storing and transporting meat and in transporting animals intended for slaughter.

The act is under the ministry responsible for veterinary services and some of the duties under it require consultation with the minister responsible for health. Moreover because it deals with meat for export, the ministry responsible for trade and industry is also a relevant player in the implementation of the act.

*Standards Act (Chapter 496), 1974*

The Standards Act is the main legislation on standards formulation and implementation in Kenya. Section 3 establishes the Kenya Bureau of Standards (KEBS), whose function, *inter alia*, is to make arrangements and provide facilities for the examination and testing of commodities and the substances used to manufacture, produce, process or treat them. In a nutshell, KEBS seeks to ensure the safety of products and ingredients.

Section 6 of the act creates the National Standards Council mandated to declare any specification or code of practice framed by KEBS to be a Kenyan standard. Members of the council include the Chair, public officers and other persons with knowledge of industrial or commercial standards (appointed by the minister responsible for trade).

The minister is empowered under the act to appoint inspectors who are mandated at all reasonable times to inspect and take samples of any commodity or material or substance being used in its manufacture, production, processing or treatment. They can also open containers.

KEBS works closely with three main public bodies in the development and implementation of health standards on animal and animal products, plant and plant products and food safety. These are KEPHIS, DVS and Ministry
of Health. KEBS is the contact point for the Codex Alimentarius Commission\(^3\) and the International Organization for Standardization (ISO).

KEBS also has technical committees to deal with different issues and to assist in the development of standards. These committees comprise 12–15 members drawn from industry, regulatory authorities, the Weights and Measures Department, consumer organizations, institutions of higher learning, research organizations and non-governmental organizations.

KEBS is a member of the National Biosafety Committee and has a memorandum of understanding with NEMA.

*Public Health Act (Chapter 242), 1921*

The Public Health Act makes provision for securing and maintaining health. The act establishes a Central Board of Health (sect. 3), which is empowered to advise the Minister of Health on all matters affecting health. It contains provisions that ensure the protection of foodstuffs intended for human consumption (sects. 127–28). Another significant provision for food safety relates to the requirement that local authorities ensure that water supplies, food and milk are in good condition.

### 2.3. Biodiversity management and biosafety

Kenya ratified the Convention on Biological Diversity (CBD)\(^4\) in 1994, which provided the impetus for the crafting of a framework environmental law to provide the normative and institutional anchorage for the conservation, sustainable use and equitable sharing of benefits emanating from biodiversity in consonance with the CBD’s objectives.

#### 2.3.1. Environmental Management and Coordination Act

The framework law, the Environmental Management and Coordination Act (EMCA), No. 8 of 1999, provides for the establishment of a legal and institutional framework for the management of the environment and for matters connected and incidental thereto. The promulgation of this act was aimed at ensuring that there was an overarching legal framework to guide environment

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\(^3\) For a discussion of Codex, see Chapter 2, Part V.

\(^4\) For a discussion of this convention, see Chapter 2, Part VI.
management in the country and to provide coordination within and among the various sectoral laws and agencies dealing with environmental matters. Section 3 of the act declares the right of every Kenyan to a clean and healthy environment as well as the corresponding duty to safeguard and enhance the environment. The entitlement to a clean and healthy environment includes access by any person in Kenya to the various public elements or segments of the environment for recreational, educational, health, spiritual and cultural purposes.

Section 5 lays out the principles of sustainable development which underpin the act. These are:

(a) the principle of public participation in the development of policies, plans and processes for the management of the environment;
(b) the cultural and social principles traditionally applied by any community in Kenya for the management of the environment or natural resources insofar as these are relevant and not repugnant to justice and morality or inconsistent with any written law;
(c) the principle of international co-operation in the management of environmental resources shared by two or more states;
(d) the principles of inter-generational and intra-generational equity;
(e) the polluter-pays principle; and
(f) the precautionary principle.

These principles undergird the attainment of Biosecurity.

Protection of the environment

EMCA provides for the protection and conservation of various ecosystems. For instance, it bars the introduction of animals whether alien or indigenous into lakes or wetlands (sect. 42(c)) or the deposit of any substance likely to have an adverse environmental effect on a river or lake (sect. 42(e)). To ensure sustainable management of such ecosystems, EMCA provides for the development of a management plan that addresses, among other issues, prevention and control of pollution, guidelines for access to and exploitation of resources and the overall management of biodiversity (sect. 42(3)).

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Section 50, dealing with the conservation of biological diversity, deserves special mention. It requires NEMA, the body charged with coordinating all matters relating to the environment in Kenya, to prohibit and control the introduction of alien species into natural habitats. It is expected that NEMA will issue guidelines on this function but this is yet to be accomplished.

NEMA is also mandated to issue guidelines for the sustainable management and utilization of genetic resources for the benefit of the people of Kenya. These include guidelines on access to genetic resources, the sharing of benefits derived from the genetic resources, biosafety measures necessary to regulate biotechnology and measures necessary to regulate the development, access to and transfer of biotechnology (sect. 53). Under this provision, NEMA has drafted two sets of regulations, some dealing with access to genetic resources and others dealing with biosafety. The former have not been promulgated while the latter informed the development of the draft Biosafety Bill which has not been finalized.

Environmental impact assessment

Environmental impact assessment (EIA) is a process which enables the examination, analysis, and assessment of proposed projects, policies or programmes and the integration of environmental issues into development planning in order to maximize the potential for environmentally sound and sustainable development. The EIA process, as argued by Hunter and others, should ensure that before granting approval (1) the appropriate government authorities have fully identified and considered the environmental effects of the proposed activities under their jurisdiction and control and (2) affected citizens have an opportunity to understand the proposed project or policy and to express their views to decision makers”.6

EMCA identifies the areas in which an EIA must be carried out, which include biosafety.

Environmental audit and monitoring

Under EMCA, NEMA is charged with the responsibility of ensuring that environmental audits are carried out for all activities likely to have significant

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effects on the environment (sect. 68). It is also mandated to monitor all environmental phenomena with a view to making an assessment of any possible changes in the environment and the possible impacts; and the operation of any industry, project or activity with a view to determining its immediate and long-term effects on the environment (sect. 69). Under this mandate, NEMA can ensure plant and animal health by protecting the integrity of the environment which comprises their habitat.

**Regulatory institutions**

To ensure conformity with its provisions, EMCA puts in place an elaborate institutional framework. The National Environment Council (NEC) is a top policy-making body under the act charged with the responsibility of formulating policy on matters relating to environment management in Kenya, setting national goals and objectives and determining policies and priorities for the protection of the environment; and promoting co-operation among public departments, local authorities, private sector actors, non-governmental organizations (NGOs) and other organizations engaged in environmental protection programmes. Those who sit on the council include two representatives of public universities in Kenya, two representatives of specialized research institutions, three representatives of the business community and two representatives of NGOs active in the environmental field. As already noted, NEMA is the principal government institution responsible for the implementation of all policies relating to the environment.

The Public Complaints Committee (PCC) is set up under Section 31 of the act. It investigates complaints relating to environmental damage and degradation generally and it has powers to investigate NEMA. It can also initiate investigations on its own without waiting for a complaint to be made. The mandate of the PCC is wide enough to cover animal and plant health. Some of the complaints that have been brought to the PCC include concerns about the introduction of alien species and their impacts on plant and animal health.

The National Environment Tribunal (NET) is established under Section 125 of EMCA. The tribunal is set up to hear appeals from administrative decisions taken by organs responsible for enforcement of the provisions of EMCA which encapsulate diverse aspects of plant and animal health.

Established under Section 70 of EMCA, the Standards and Enforcement Review Committee is required, among other things, to advise NEMA on
water quality procedures and standards and discharge of effluents into the environment; air quality and emission standards; standards for waste disposal and management, hazardous waste, pesticides and toxic substances. These standards have direct and indirect implications for plant and animal health as well as food safety.

**Compliance and enforcement mechanisms**

EMCA contains provisions for compliance and enforcement, thus contributing towards the achievement of *Biosecurity*. These mechanisms include:

1. Environmental restoration orders requiring restorative action, preventing harm to the environment, payment of compensation for harm to the environment and levying charges commensurate with costs of restoring degraded environments (sects. 108–111).
2. Environmental easements for facilitating the conservation and enhancement of the environment through the imposition of obligations in the form of environmental conservation orders in respect of the use of land (preservation of flora and fauna; preservation of water flow, open space, scenic view among others) (sects. 112–115).
3. Fiscal incentives and disincentives to induce or promote proper management of the environment. These include customs and excise waivers, tax rebates, tax disincentives to deter bad environmental behaviour and user fees.
4. Criminal sanctions.

**2.3.2. Biosafety legislation**

The Cartagena Protocol on Biosafety is a protocol to the CBD which has been ratified by Kenya. It had already become apparent during the negotiations for the CBD that further work was required towards a biosafety protocol.

Prior to the ratification of the protocol, Kenya had developed *Draft Regulations and Guidelines for Biosafety in Biotechnology for Kenya* in 1998 under the UNEP-GEF Pilot Project on the development of national biosafety frameworks. These were issued by the National Council for Science and

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7 For a discussion of this instrument, see Chapter 2, Part VII.
Technology (NCST) established under the Science and Technology Act, Chapter 250 (1977). They are the main instrument for regulating GMOs in Kenya to date and require that the release of GMOs be preceded by the approval of the National Biosafety Committee (NBC). Membership in the NBC includes representatives of relevant institutions (KEPHIS, NEMA, KEBS, DVS) and line ministries (environment, health, agriculture, education, science and technology). The Office of the Attorney-General is also represented, to provide advice on relevant emerging legal issues. The relevant regulatory authorities must undertake risk assessments, and thus require information from applicants such as a description of the GMOs and their intended uses in Kenya.

For crops, KEPHIS is the relevant authority, advising the NBC on whether or not to allow imports and what to do after the assessment. The guidelines provide that it is an offence to import GMOs without prior approval of the NBC. Penalties for offences under the biosafety regulations were left to be made by the minister. In order to do this the minister requires powers conferred upon him or her by an act of parliament. To date, this has not been done although there are some prescribed penalties in draft form under the proposed National Biosafety Bill.

The main aim of the regulations is to enhance effectiveness in the use of new products and to ensure safety regarding human health and the environment. They require institutions carrying out work on genetic modification to establish institutional biosafety committees. These committees are required to advise their respective institutions on drawing up proposals that take cognizance of applicable biosafety measures and advise them on activities that should be brought to the attention of the NBC. The applicant is expected to make an application in the prescribed form detailing all information on the proposed work and send the form to the NCST as the secretariat of the NBC. The NBC should acknowledge receipt within 30 days and verify the information for completeness using a checklist, whereupon it may request additional information from the applicant within 60 days. Deliberation must be within 90 days and the decision to approve or deny approval communicated within 210 days.

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8 The proposed penalties may not achieve the desired goals as they are relatively lenient. For example, someone who imports GMOs without the advance informed agreement of the country of import may only be liable to a fine not exceeding fifty thousand shillings (about €530 at today’s prices). Under such circumstances potential violators may find it convenient to commit the offence and pay the fine.
The NBC gives approvals for GMO events while KEPHIS gives permits where plant materials are involved. The NBC approval is predicated on the applicant satisfying phytosanitary conditions and getting a permit from KEPHIS. KEPHIS appoints technical experts to coordinate risk assessments on behalf of the NBC where plant materials are at issue.

There is also close collaboration between VSD and NBC. For instance, the application by a private company to use GM soya for piglets was referred to the NBC by VSD. Similarly, the PCPB referred to the NBC an application made to it regarding a GM biopesticide for use in rose farming.

The draft National Biosafety Bill is an attempt to give a firm legal basis to biosafety regulation in Kenya. It seeks to align the draft regulations with the Cartagena Protocol. The main objective of the draft biosafety bill, issued by the NCST under the Science and Technology Act within the auspices of the Ministry of Education, Science and Technology, is to ensure an adequate level of protection in the field of safe transfer, handling and use of GMOs resulting from modern biotechnology that may have an adverse effect on the environment; and to establish a transparent and predictable process for review and decision-making on such GMOs and related activities (sect. 4).

Section 5 establishes the National Biosafety Authority, the board of which comprises:

(a) a Chairperson, who shall be an eminent scientist, appointed by the minister responsible for science and technology matters;
(b) three other members comprising experts in biological, environmental and social sciences;
(c) the Permanent Secretaries in the ministry responsible for science and technology and the Ministry of Finance;
(d) the Director-General of NEMA;
(e) the Managing Director of KEBS;
(f) the Managing Director of KEPHIS;
(g) the Director of VSD;
(h) the Secretary to the National Council for Science;
(i) the Chief Public Health Officer;
(j) the Director of Agriculture; and
(k) a representative of the consumer information network.
The functions of the authority as enumerated under Section 7 include to:

(a) receive, respond to and make decisions on applications under the bill;
(b) establish administrative mechanisms to ensure the appropriate handling and storage of documents and data in connection with the processing of applications and other matters covered by the bill;
(c) establish contact and maintain liaison with other countries and organizations dealing with biosafety;
(d) establish a database for the purpose of facilitating collection and dissemination of information relevant to biosafety;
(e) identify national requirements for manpower development and capacity building in biosafety;
(f) maintain a directory of experts in biotechnology and biosafety;
(g) keep a record of biotechnology and biosafety activities in Kenya;
(h) advise institutions and persons on mitigation measures to be undertaken in case of accident; and
(i) promote awareness and education among the general public in matters relating to biosafety.

The authority is further granted powers to approve or reject applications as well as to determine whether or not to carry out risk assessments. The following are activities subjected to the written approval of the authority:

(1) contained use involving GMOs;
(2) introduction of GMOs into the environment;
(3) importation and placing of GMOs into the market; and
(4) transportation of GMOs through Kenya.

Any decision made by the authority is subject to review upon the request of a regulatory body or any applicant in situations where new scientific information relating to biosafety of the GMOs is discovered or there has been a change of circumstances. Any applicant who, having knowledge of such information, withholds it from the authority, commits an offence under the bill. Furthermore, failure to adhere to any of the requirements on approvals constitutes an offence.

Any person aggrieved by any decision made by the authority has the right of appeal to an Appeals Board established under Section 26 of the bill. It is also noteworthy that the bill identifies regulatory agencies responsible for different issues. These are contained in the first schedule and include the
Ministry of Health, VSD, KEBS, KEPHIS, the Kenya Industrial Property Institute, the Kenya Wildlife Service, PCPB and NEMA. They are in charge of the following:

- monitoring of applicants' activities to ensure that they conform to the bill;
- informing the authority of any new information aimed at enhancing the continued safe use of GMOs; and
- inspecting and evaluating activities involving GMOs.

Biosafety inspectors are appointed by the minister responsible for science and technology, and are accorded a variety of powers. In addition to criminal sanctions, the draft bill anticipates the use of other means of redress or damage resulting from GMOs. It provides at Section 42 that "liability and redress for any damage that occurs, as a result of activities subject to this Act, shall be addressed by applicable laws". There is concern, however, that this provision is not adequate because the applicable laws predate biotechnology activities and may not cover all kinds of damage likely to arise from them.9

III. ANALYSIS OF ADEQUACY OF BIOSECURITY LEGAL FRAMEWORK

3.1. Institutional fragmentation and conflicting mandates

It is clear from the above analysis that Kenya has many laws dealing with Biosecurity. The challenge of implementation and coordination of the various laws however remains. At both the normative and institutional level, there are overlaps which create room for conflict. Additionally, many of these institutions may perform their duties within narrow confines and thus fail to consider national imperatives for cohesion and synergies.

From the analysis above, it is clear that the institutional basis for Biosecurity is dispersed among different ministries and institutions. These include:

1. Ministry of Agriculture
   - KEPHIS
   - PCPB

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2. Ministry of Livestock Development and Fisheries
   - VSD
   - Department of Fisheries
3. Ministry of Health
   - Department of Public Health
   - Public Health (Standards) Board Central Board of Health
   - Central Board of Health
   - Medical Department
4. Ministry of Education, Science and Technology
   - NCST
   - NBC
5. Ministry of Local Government
   - Local authorities
6. Ministry of Trade and Industry (MOTI)
   - KEBS
   - National Committee on the World Trade Organization (WTO)
7. Ministry of Environment
   - NEMA
   - NEC
   - NET
   - PCC
8. Ministry of Finance
   - Customs and Excise Department, working with KEPHIS and KEBS at points of entry
9. Ministry of Justice
10. Ministry of Tourism.

To bring together the institutions responsible for different regulatory functions, a number of inter-ministerial coordinating committees have been established. For instance, there is an inter-ministerial committee to advise the government on all matters pertaining to the WTO. This committee later metamorphosed into the National Committee on WTO which includes governmental as well as non-governmental actors. It includes the Attorney-General, the Office of the President, MOTI, and the Ministries of Finance, Planning and National Development, Health, Agriculture, Foreign Affairs, Labour and Human Resources, Environment and Natural Resources, Information and Communications and Transport. The ministries act as the focal points for sub-committees handling relevant WTO issues and each line
ministry handles its core issues. KEBS, KEPHIS and NEMA are also represented on the committee. The coordinating ministry is MOTI.

On a positive note, the establishment of KEPHIS in 1996 has led to greater coordination of the phytosanitary aspects of Biosecurity. This has however been hampered by the delay in amending the relevant laws to entrench the role of KEPHIS. It is particularly a matter of concern that the proposed bill to institutionalize KEPHIS is yet to be promulgated and continues to be debated.10 Given the centrality of KEPHIS in the Biosecurity framework in Kenya, its lack of a definitive status, being an institution established only through a ministerial order that, at least in theory, could be revoked creates uncertainty that does not augur well for Biosecurity.

The Plant Protection Act, Suppression of Noxious Weeds Act, Seeds and Plant Varieties Act and Agriculture Act all need to be amended to synchronize their provisions with the role of KEPHIS. Currently, there is room for conflict between the Ministry of Agriculture officers and KEPHIS officers in the performance of their duties. One example is where a Seed Committee chaired by the Agriculture Secretary and comprising seed sector stakeholders made a decision which was against the Seeds and Plant Varieties Act yet KEPHIS was expected to implement it.

Another positive trend is the involvement of officers from line institutions in related regulatory bodies such as the involvement of KEPHIS, VSD and KEBS in NEMA, the NBC and the National Committee on WTO. This has assisted in coordination of various Biosecurity functions.

There is a need to examine the legal status of the various institutions involved in Biosecurity. VSD remains a department of the government while KEPHIS is a state corporation with more autonomy and flexibility. This perhaps reflects the emphasis the government places on agriculture. Given the constant change in line ministries where there is sometimes a livestock ministry and other times the ministry’s functions are brought together with agriculture, it is important to create VSD as an autonomous body like KEPHIS. It could for instance be brought under an SPS regulatory regime that has divisions dedicated to the roles that KEPHIS and VSD currently play. Proposals to establish one such agency, the Kenya Animal and Plant

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10 A version of the bill was tabled for discussion at a meeting on 3 May 2007, four years since the last draft was finalized and presented to the ministry.
Health Inspectorate Service (KAPHIS) to encompass both KEPHIS and VSD, have been made over the years but they are yet to materialize.

The food safety aspect of Biosecurity also needs to be streamlined. For instance, the biosafety regime needs to be harmonized with the Food, Drugs and Chemical Substances Act, the Standards Act and the Public Health Act. The roles of the Ministry of Health, the Ministry of Agriculture, KEBS and VSD need to be synchronized and rationalized. This need has been recognized and a Food Safety Committee set up with the Agriculture Secretary as Chair, to provide guidance on synergies. This committee is the focal point for all food safety issues and draws its membership from the Ministries of Agriculture, Health and Trade, the Kenya Medical Research Institute, KEBS and the Kenya Agricultural Research Institute. It was launched on 4 May 2007.

This raises a broader issue of the diverse inspection functions. Under most of the laws discussed above, there is provision for inspectors. This function needs to be coordinated so that training and the inspection function are carried out in a systematic way. Currently, inspectors are few and this hampers the effective discharge of the duties entrusted to the officers. For instance while KEPHIS is required to provide border control with respect to plant materials (a function previously performed by airport staff of the Ministry of Agriculture), there are not enough inspectors to cover all entry points. Moreover, there are no VSD inspectors to address meat and meat products at the points of entry.

Another challenge for Biosecurity is the operation on the basis of drafts with regard to biosafety. Kenya has ongoing work on GMOs, but the permits have been issued only on the basis of draft regulations, as noted above. The process of promulgating a biosafety law has remained protracted. Attempts to come up with a National Biosafety Act in 1999 failed. There is currently, as was pointed out above, a draft Biosafety Bill which has been the subject of discussion since 2002.

It is worth noting that Kenya also has a biotechnology policy promulgated in December 2006 which seeks to provide a framework for safe development and application of biotechnology. There is also an ongoing process to develop a national biotechnology strategy. The policy framework needs to be supported by a law and a national strategy to mainstream the application of biotechnology in national development. In the absence of a finalized law, the
NBC continues to consider applications and allow work on GMOs on the basis of the draft regulations.

As Kenya refines its *Biosecurity* framework, it has to contend with the imperatives of regional integration. Membership in sub-regional and regional groupings will have implications for the efficacy of the framework put in place. Kenya is a member of the East African community, one of whose objectives is harmonization of laws and standards to facilitate the creation of a common market. This has implications for the national *Biosecurity* framework. Similarly, membership to the Association for Agricultural Research in Eastern and Central Africa, which is currently working on harmonizing seed and biosafety laws, will have implications for Kenya’s national framework and institutions.

### 3.2. Proposals for reform

It is widely recognized that there is need for greater institutional synergy. The historical separation of SPS functions calls for sensitization of stakeholders and consensus building on the benefits of the establishment of a single agricultural regulatory authority, which could also address *Biosecurity*. The draft Agriculture Sector Reform Bill, 2006, sets out to reorganize and update the legal and regulatory environment in the agricultural sector by consolidating roles in three proposed agencies:

1. the Agriculture Development Board;
2. the Livestock Development Board; and
3. the Agriculture Sector Regulatory Authority (ASRA).

The proposed ASRA draws its membership from a wide array of actors including the Ministries of Livestock and Fisheries, Agriculture, Health and Treasury, the universities, the Agriculture Development and Livestock Development Boards, the Kenya Association of Manufacturers and two representatives from the agribusiness sector.

The objective of the bill is laudable and its timeliness cannot be gainsaid. Currently, the agricultural sector is governed by over 130 pieces of legislation, 60 of which regulate and control various commodity sectors. Many of these laws are outdated and in need of repeal or amendment. The consolidation of legislation is seen as a way of fast-tracking and updating laws in a context where legislation takes a long time to go through
parliament. The Law Reform Commission, the national body charged with reviewing, updating and drafting new laws, is currently developing many bills. It has been historically inactive but has in the last five years developed a strategy for law reform. The demand for new laws far outstrips the capacity of the commission and drafters in the Attorney-General’s chambers. Consequently, many draft laws are being generated by the sector that still require amendment, review or overhaul.

This legislative initiative presents an opportunity to mainstream Biosecurity and to provide a coordinated institutional framework for its implementation. There is no reference to Biosecurity in the Strategy for Revitalizing Agriculture (SRA) 2004–2014 concluded in 2005, although it can certainly be read to embrace Biosecurity. The SRA identifies six interventions to reverse the decline in the agriculture sector and position it competitively in the global arena:

(a) review and harmonization of the legal, regulatory and institutional framework;
(b) restructuring and privatization of non-core functions of parastatals and ministries;
(c) improving the delivery of research, extension and advisory support services;
(d) improving access to quality inputs and financial services;
(e) improving access to both domestic and external markets; and
(f) formulating food security policy and programmes.

These interventions have direct and indirect implications for Biosecurity. Part 2.3.4. on animal health and plant protection services is of direct relevance. It seeks to revamp animal health and plant protection services in order to increase production through:

(a) reviewing and enforcing laws on delivery of animal health, fish and plant protection services;
(b) building capacity for laboratory analysis for diagnosis and remedial action;
(c) ensuring development and maintenance of infrastructure for the control of livestock and plant diseases; and

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11 Cabinet memo on Consolidating Legislation in the Agricultural Sector, 2007.
(d) enforcing sanitary, phytosanitary and zoosanitary measures to prevent introduction and spread of new pests and diseases.

The opportunity to include Biosecurity is there in the anticipated changes but it requires clarification and articulation. First, it is not clear from the draft Agriculture Sector Reform Bill how the existing regulatory institutions will interface with the proposed ones. Will they be absorbed in the ASRA? Second, it is not clear how the authority will be structured to perform all the regulatory roles currently dispersed in different agencies. Third, it is not clear how the regulatory roles will be linked to regulatory agencies in other sectors that have relevance for the agriculture sector. Failure to clarify and articulate synergies will confound an already complicated scenario.

IV. CONCLUSION AND RECOMMENDATIONS

Kenya has in place an elaborate Biosecurity legal and institutional framework. This framework has been refined over time and continues to evolve. There is recognition of the need to amend laws to ingrain elements of Biosecurity from the agriculture, health and environmental angles. While there is a framework environmental law, the drafting of the implementing regulations of that law is ongoing. For instance, it is expected that regulations on alien and invasive species will be put in place. This chapter has captured the framework as it grapples with these changes.

The implementation of the framework is a challenge. It was not easy to ascertain, in carrying out the present review, the extent of conformity with existing regulations. Furthermore, since most of the laws and institutions have functions other than Biosecurity, it was not possible to rate the effectiveness of the normative and institutional frameworks. For instance, the emphasis on trade of agricultural products and exports of meats seems in some cases to override concerns for national Biosecurity. In addition, the emphasis on certain kinds of products leaving out others can compromise Biosecurity. There is, for instance, evidence that game meat consumption has gone up in Kenya in the last ten years, but because dealing with game products is illegal, most of such meat finds its way to the marketplace without the requisite inspection. This is an example of the difficult context that can compromise the implementation of even the best normative and institutional framework for Biosecurity.
At another level, the framework still has normative and institutional overlaps as new laws and institutions are put in place. The challenge is to ensure that these laws and institutions seek to achieve the same objectives. There are many international instruments which Kenya has ratified and whose implementation is in the nascent stages, and the challenge of synergizing these *inter se* as well as with national development imperatives will continue. Membership in international, regional and sub-regional bodies will also influence the evolution of the national *Biosecurity* framework.

Capacity to implement an effective national *Biosecurity* framework is another issue of concern. It is not helped by the overlaps which can lead to bureaucratic delays as each body looks to the other to act. It could also drain necessary capacity from implementation which instead must deal with conflicts. While it may not be possible to contain all elements of *Biosecurity* in one law or institution, there is a need for a focal point to ensure that the overall objectives are not compromised by actions within any one function in the framework. Such a focal institution would be the repository of information as well as the mechanism for dissemination of that information to all relevant actors. It needs to be able to oversee the entire framework and have the authority to require certain actions on the part of all actors to ensure more synergy and cooperation.

At a broader level, inter-institutional rivalries have hampered the development of a coherent *Biosecurity* normative and institutional framework. This raises the need for awareness raising among stakeholders and engagement in consensus building to ensure buy-in by all stakeholders. The SRA and accompanying reforms provide an opportunity for consensus building and banking on political goodwill. It is worth noting that within this framework, an inter-ministerial Cabinet committee has been established to work with the Agriculture Sector Coordinating Unit and technical committees to fast-track the thematic interventions identified under the SRA.
UGANDA COUNTRY STUDY*

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* This chapter was prepared by Judy Obitre-Gama.
I. INTRODUCTION

Biosecurity measures in agriculture are needed to protect agricultural production systems and those dependent on these systems. Producers and others dependent on agriculture can see their livelihood destroyed by animal and plant pests and diseases or damage to the environment such as impacts resulting from invasive alien species. Measures are needed to protect human health, particularly of vulnerable groups that can be exposed to severe health risks, which Biosecurity attempts to prevent. Biosecurity seeks to protect the environment, promote sustainable production and build consumers’ confidence in agricultural products. Public awareness of environmental issues and human dependency on biodiversity have resulted in numerous commitments to achieving sustainable development, and achieving these will require an effective approach to Biosecurity.

This report appraises legislation and policy in Uganda on Biosecurity comparing the national framework with the international regime on Biosecurity. It concludes by proposing a way forward for an effective Biosecurity legal regime in Uganda.

II. NATIONAL LEGAL REGIME ON BIOSECURITY

2.1. Constitution

Constitution of the Republic of Uganda, 1995

The Constitution, as amended in February 2006, in the National Objectives and Directive Principles of State Policy, provides that the state shall protect important natural resources including fauna and flora on behalf of the people of Uganda (Objective XIII). The state commits itself to promote sustainable development and the rational use of natural resources so as to safeguard and protect the biodiversity of Uganda (Objective XXVII).

The right to a clean and healthy environment is enshrined in article 39 while article 245 requires parliament to pass laws for the protection and preservation of the environment.
2.2. Environment and wildlife

National Environment Act (Chapter 153), 1995

The objective of the National Environment Act (NEA) is to provide for sustainable management of the environment and to establish an authority as a coordinating, monitoring and supervisory body for that purpose.

The National Environment Management Authority (NEMA) is established under the act as the principal agency in Uganda responsible for the environment. NEMA is required to ensure the integration of environmental concerns into overall national environmental planning through coordination with the relevant ministries, departments and agencies of government; and initiate legislative proposals, standards and guidelines on the environment in accordance with the act. NEMA is mandated to ensure the observance of proper safeguards in the planning and execution of all development projects, including those already in existence that have or are likely to have a significant impact on the environment. In a similar vein, NEMA shall review and approve any environment impact assessment (EIA) or statement submitted in accordance with the laws of Uganda.

Accordingly, an EIA shall be undertaken by the developer where the lead agency, in consultation with NEMA, is of the view that the project proposed may have an impact, is likely to have a significant impact or will have a significant impact on the environment. The list of projects that must have an EIA is contained in the third schedule to the act. The list includes the introduction of new crops and animals (item 8), and the introduction of fauna and flora into ecosystems of natural conservation areas (item 13). The minister on the advice of the board of directors of NEMA may amend the schedule.

NEMA, in consultation with the lead agencies, is enjoined to take all measures to ensure that biodiversity is conserved in situ, where possible, and ex situ where not. Further provision for preservation of wildlife in situ is made in the Uganda Wildlife Act (see below). The NEA stipulates that access to genetic resources shall be regulated in order to sustainably utilize genetic resources for the benefit of the people of Uganda.
National Environment
(Access to Genetic Resources and Benefit Sharing) Regulations, 2005

The Access to Genetic Resources and Benefit Sharing Regulations define genetic resources as genetic material of actual or potential use or value and includes their derivative products and intangible components, while "access" is defined to mean the obtaining, possessing and using of genetic resources for purposes of research, bio-prospecting, conservation, industrial application or commercial use. The regulations prescribe the procedures for access for these purposes.

The regulations apply to access to genetic resources or parts thereof, whether naturally occurring or naturalized, whether in in situ conditions or ex situ conditions, including genetic resources bred for or intended for commercial purposes within Uganda or for export. Excluded from the application of the regulations are the exchange of genetic resources that are certified to be purely for food or other consumptive purposes as prescribed by the relevant laws; the transit of genetic resources through Uganda; exchange by a local community inter se for consumptive purposes; and access to genetic resources derived from plant breeders as defined by laws relating to plant breeding and plant varieties. Where access to genetic resources is likely to have a significant impact on the environment, an EIA shall be carried out prior to the conclusion of a material transfer agreement.

The Uganda National Council for Science and Technology (UNCST), which is established under the Uganda National Council for Science and Technology Act, Chapter 209, 1990 (see below), is designated as the competent authority for the purpose of fulfilling the object of the regulations. UNCST is enjoined to coordinate all activities of lead agencies relating to access to genetic resources in accordance with these regulations and the NEA; and submit to NEMA reports relating to the implementation of these regulations. NEMA, however, retains the function of initiating the formulation of national policy on access to genetic resources; developing guidelines for access to and export of genetic resources; collaborating with lead agencies in carrying out public awareness campaigns, designing capacity building programmes; ensuring compliance with and enforcement of these regulations; developing guidelines for the export of genetic resources and benefit sharing; and advising on access to genetic resources outside protected areas.
Uganda Wildlife Act (Chapter 200), 2000

The Uganda Wildlife Act provides for the sustainable management of wildlife which is defined to include any wild plant of a species native to Uganda. Any species of plant whose natural range does not now or did not in the past include a specific part of Uganda or the whole of Uganda is referred to as an alien species (sect. 1) and the act does not authorize the introduction of alien species of plants or animals into wild habitats within Uganda (sect. 2). Though a wild habitat is not specifically defined, it is construed to refer to a wildlife conservation area declared as such by statutory instrument (sects. 17 and 18).

This act is intended to promote the conservation of wildlife in Uganda in order to maintain the biological diversity that exists for the benefit of the people of Uganda. This includes the protection of rare, endangered and endemic species of wild plants (sect. 2). The minister responsible for wildlife may, by statutory order, declare any species of wild plant or wild animal as a protected species under the act (sect. 27). Use of resources in wildlife protected areas is permitted by the Uganda Wildlife Authority (UWA) in instances where a permit is issued to an applicant specifying the extent and duration under which such applicant has access to the resources stated in the permit (sect. 23).

The UWA is established under the act to, inter alia, ensure the sustainable management of wildlife conservation areas; develop and implement policies in the field of wildlife management; establish policies and procedures for sustainable utilization of wildlife by and for the benefit of the communities living in proximity to wildlife; control the development of tourist facilities in wildlife protected areas; and promote the conservation of biological diversity ex situ and contribute to the establishment of standards and regulations for that purpose.

UWA is authorized to delegate any of its functions in writing to a lead agency, a committee or any public officer. In performing its functions, UWA shall coordinate with lead agencies involved in the field of wildlife management. A lead agency includes any ministry, department, parastatal agency or public officer in whom the law vests functions related to the management of wildlife or wildlife conservation areas and includes a local government council.
2.3. Plants and seeds

Plant Protection Act (Chapter 31), 1976

The Plant Protection Act was originally passed as an ordinance in 1937. The scope of the act can be gleaned from the long title which limits the act to the prevention of the introduction and spread of diseases destructive to plants. The definition section is very limited and is an indicator of the purpose of the act. The act precedes the International Plant Protection Convention (IPPC)\(^1\) of 1951 so arguably it does not take any international considerations into account. The administrative structure was limited to the size of the service required. Over time, the structure became too small and the budget too restrictive to allow the operation of an efficient and effective service. The penalties imposed were found to have no deterrent or actual value. For these reasons, inter alia, the act was reviewed in 2001 and the Plant Protection and Health Bill, 2003, was drafted.

The 2003 bill attempted to fill in the gaps identified in the Plant Protection Act by establishing a Technical Committee to assist the commissioner and the minister in carrying out the functions outlined for the Department of Crop Protection in the ministry responsible for agriculture. The penalties were reviewed and currency points introduced to make the penalties more realistic. The definition section was expanded to include newer terms used in phytosanitary service, drawing on the Glossary of Phytosanitary Terms of the new revised text of the IPPC (1997). The thrust of the bill was to protect plant health and the natural environment and comply with international standards on plant protection in order to enhance the international reputation of Ugandan agricultural products, especially exports. The cost recovery proposed in the draft bill particularly to enable rapid response to epidemics of quarantine importance was not included in the final text of the bill. The 2003 bill was found lacking in these respects and a revised bill was proposed in 2005.

The Plant Protection Bill, 2005, seeks to consolidate and reform the law relating to protection of plants against pests; to prevent the introduction and spread of pests that may adversely affect Uganda’s agriculture, the natural environment and the livelihood of the people; to ensure sustainable plant and environmental protection; and to regulate the export and import of plants and plant products and the introduction of new plants in accordance with

\(^1\) For a discussion of this convention, see Chapter 2, Part III.
international commitments on plant protection. The bill proposes a cost recovery mechanism to enable rapid response to epidemics of quarantine importance. It introduces pest risk analysis and strengthens the import and export controls of plants, plant products and regulated articles. The objective of the bill is to protect and enhance the international reputation of Ugandan agricultural products, and to entrust all plant protection regulatory functions to the government through the national plant protection organization (NPPO).

The department responsible for plant protection within the Ministry of Agriculture is designated as the NPPO and is responsible for the protection of the plant resources of Uganda from pests that exist in the country or could be introduced into the country. The NPPO is responsible for the implementation of the act. To this end, the NPPO is responsible for carrying out surveillance of growing plants, including areas under cultivation and wild flora, and of plants and plant products in storage or in transport, for the purpose of reporting the occurrence, outbreak and spread of pests, and of controlling those pests.

The NPPO is mandated to enforce the act and any other legislation relating to plant protection that the minister may identify; and to establish procedures for accreditation of any quarantine station, official analyst, official laboratory or any other person or institution from the public or private sector involved in phytosanitary matters. The minister is to appoint a Plant Protection Technical Committee to advise the commissioner on all technical matters arising from the administration of the act and on any other related issues. The commissioner shall be the head of the NPPO and responsible for the day-to-day administration of the act. The commissioner is defined to mean the commissioner responsible for plant protection or any other commissioner or competent person assigned by law to administer the act.

The minister is authorised, from time to time, to appoint by notice in the gazette, officers of the NPPO or other competent persons to be inspectors for the purposes of the act. In addition, the minister has powers, by statutory instrument, to prescribe functions under the act that may be delegated to any specified competent individual or institution, including designation of laboratories and competent scientists. Delegated individuals or institutions shall be required to comply with instructions that may from time to time be issued by the minister; report on their activities to the NPPO on a periodic basis as may be determined by the minister; and assist in and cooperate with the NPPO in attaining the purposes of the act.
A perusal of related laws reveals that there are overlaps that should be addressed in the bill to avoid institutional conflict and the resultant inefficiency it engenders. Section 12 of the Agricultural Seeds and Plant Act (Chapter 28, 1994), authorizes the National Seed Certification Service (NSCS) to establish phytosanitary standards and practices for any particular crop as the need arises. The NSCS is further authorized to direct that seeds or plants harbouring pests and diseases be destroyed within a specified period of time and in a specified manner. The Cotton Development Act (Chapter 30, 1994), in Section 12, mandates the minister responsible for agriculture, in consultation with the Cotton Development Organization, to direct that any cotton seed or plant harbouring or likely to harbour any cotton pest or cotton disease be destroyed. A provision is proposed in the Plant Health and Protection Bill, 2005, to address this anomaly and ensure that the principal authority for all phytosanitary matters is assigned by law to the NPPO.

National Forestry and Tree Planting Act, 2003

The National Forestry and Tree Planting (NFTP) Act contains provisions that are contrary to the purpose of establishing an NPPO with principal responsibility for phytosanitary services in Uganda. Under Section 36 of the NFTP Act the minister responsible for forestry, the National Forestry Authority or a district council is authorized to notify the public through the mass media of the existence of plant and livestock pests or diseases dangerous to forests or forest produce and prescribe the measures to be taken to control or eradicate those pests and diseases. Section 92(2)(g)–(i) empowers the minister by statutory instrument to issue regulations that may provide for the notification of plant and livestock pests and diseases dangerous to forests and forest produce and the measures to be taken to control or eradicate the notified pests or diseases; and the introduction of alien and exotic species.

While the authority to notify the public of plant pests or diseases as stipulated in the NFTP Act may not in itself be inconsistent with the revised text of the Plant Protection Bill, particularly if that notification enables the NPPO to improve in the efficient delivery of service, the prescription of control and eradication measures, and the authorization for the introduction of alien or exotic species (particularly as pertains to the phytosanitary health and safety of such species) are matters that are within the exclusive domain of the NPPO. To solve any mischief that may be occasioned by the aforementioned sections of the NFTP Act, a new clause was introduced in the
revised bill that stipulates that any law existing immediately before the coming into force of this act relating to plant protection shall have effect subject to such modifications as may be necessary to give effect to the bill; and where any such law conflicts with the bill, the provisions of the bill shall prevail. It is hoped that this clause will preserve the role of the NPPO as the government agency charged with the responsibility for all phytosanitary matters.

Agricultural Seeds and Plant Act (Chapter 28), 1994

The Agricultural Seeds and Plant Act provides for the promotion, regulation and control of plant breeding and variety release; multiplication, conditioning, marketing, importing and quality assurance of seeds and other planting materials. The National Seed Industry Authority established under the act is responsible for advising government on national seeds policy; constantly reviewing the national seed supply; and coordinating and monitoring the public and private seed sector in order to achieve the national seed programme objectives (sect. 3). The NSCS is responsible for the design, establishment and enforcement of certification standards, methods and procedures in the seed industry (sect. 6); while the variety release committee reviews and maintains the national variety list including the approval of new varieties of seeds, and approves variety release and entry of seeds into the seed multiplication programme (sect. 5). All biosafety issues are referred to and handled by UNCST in accordance with the law.

The Seed and Plant Bill, which was assented to and should be gazetted as an act of parliament shortly, will repeal Chapter 28. The bill establishes a National Seed Board (NSB), with the department of Crop Protection providing the secretariat. The Variety Release Committee is maintained under the bill. Seed import permits shall be issued by the NSB. Risk assessment in terms of plant health is done by phytosanitary services. The seed import permit constitutes a "no objection" to import seeds, subject to phytosanitary measures.

2.4. Animals

Animal Diseases Act (Chapter 38), 1964

The Animal Diseases Act defines animals that are within its ambit to mean all stock, camels and other ruminating animals, cats and dogs (sect. 1(a)); and disease to mean any disease contained in the list under Section 1(d). All
diseased animals or animals suspected to be infected by disease must be separated from the other animals by the owner or caretaker of the animals; and a veterinary officer or administrative officer should be notified accordingly (sect. 2). The veterinary officer notified must, once he or she has ascertained the existence and nature of the disease, report the matter to the Commissioner of Livestock and Entomology (sect. 3). An administrative officer shall, on being satisfied as to the existence of a disease affecting stock (cattle, sheep, goats, horses, mules, donkeys and poultry) within his or her area of jurisdiction, cause all owners and occupiers of farms and owners of stock in the neighbourhood to be notified of the disease (sect. 4).

All diseased or suspect animals or any animal which has been in contact with a diseased animal or has been exposed to the infection or contagion of disease shall be slaughtered on the instruction of the veterinary officer or administrative officer and the carcass disposed of according to such instruction.

**Cattle Traders Act (Chapter 43), 1964**

The Cattle Traders Act provides for the regulation of cattle trading in Uganda which can only be undertaken once a licence in the prescribed form has been issued by a veterinary officer indicating the area(s) of operation of the cattle trader (sect. 2). The licence shall be valid up to 31 December (sect. 7) and shall not be renewed if the applicant has, inter alia, been convicted of an offence under the Animal Diseases Act. A cattle trader is defined as any person who trades in cattle for the purposes of resale or slaughter (sect. 1).

**Animal Breeding Act, 2001**

The Animal Breeding Act provides for the promotion, regulation and control, marketing, import and export and quality assurance of animal and fish genetic materials. It makes general provision for the implementation of the national breeding policy in Uganda and other matters connected therewith.

The Director of Animal Resources is charged with various functions. Under Section 4, the director should promote optimum animal genetic resource management, conservation and sustainable use commensurate with Uganda’s needs and environmental protection. The Commissioner of Animal Production and Marketing is responsible for registration of animal genetic resources and related activities (sect. 6(1)), while the Commissioner of Fisheries Resources is responsible for the register of fish breeding and
related activities (sect. 6(4)). "Animals" are defined to mean all livestock, camels, donkeys, rabbits, poultry, other ruminating and pseudo-ruminating animals, fish and any other animal that the minister may by statutory instrument so declare (sect. 3).

No imports or exports of animal breeds and genetic material shall be done without obtaining a permit from the Commissioner of Livestock and Entomology (sect. 7). A list of suitable breeds for widespread use is contained in the third schedule to the act, and any breed not appearing in that schedule shall only be allowed into the country for restricted use on designated locations and experimental stations or specialized production units as approved by the director (sect. 8(1)). The director shall sanction imports and exports based on verified documentary evidence of the material being free of the disease agents and prohibited hereditary defects specified in the fourth schedule to the act (sect. 8(4)). A sample of all genetic materials defined to be semen, ova, eggs and embryos shall be submitted to a national depository for examination and future reference (sect. 9). The National Animal Genetic Resources Centre and Data Bank established under Section 13 is identified to serve as a national gene depository and examination centre for genetic materials (sect. 15(2)), among other functions. All genetic materials must conform to the national biosafety standards set by UNCST and UNBS (sect. 9).

2.5. Food

Food and Drugs Act (Chapter 278), 1964

The Food and Drugs Act makes provision for the prevention of adulteration of food which is defined to include drink, chewing gum and other products of like use or nature, and articles and substances used as ingredients in the preparation of food or drink or of such products. It excludes water, live animals or birds, animal fodder or feed and substances used only as drugs (sect. 1). The act prohibits the use of an ingredient that renders food injurious to human health in the preparation of food sold for human consumption (sect. 2) and prohibits false labelling or advertisement of food (sect. 5). Food in course of transit may be examined by an authorized officer (sect. 9).

An authorized officer means a person authorized by the Minister of Health, or a local authority with the approval of the minister. For the purposes of taking samples, an authorized person includes a police officer of or above
the rank of inspector authorized to take samples. A veterinary surgeon registered under the Veterinary Surgeons Act, in the service of the government or of a local authority, is deemed to be an authorized officer for the purposes of the inspection of animals intended for slaughter and the examination and the seizure of meat unfit for human consumption. A medical officer, a health inspector or a person having such qualifications as may be prescribed may undertake functions similar to the veterinary surgeon.

A committee known as the Food Hygiene Advisory Committee is established under the act to advise the minister on any questions relating to the act that the minister may refer to it for its consideration. The committee is appointed by the minister to hold office for such period as may be stipulated. The membership of the committee includes persons qualified to represent the interests of the public generally in relation to matters of food hygiene and related matters; and representatives of persons carrying on any trade or business affected by the operation of the act. Based on the draft Food and Nutrition Policy, the Food and Drugs Act is in the process of review through a participatory process with the involvement of the ministry responsible for agriculture in order to expand its scope from matters concerning human health to the whole food chain.

Public Health Act (Chapter 281), 1964

According to the Public Health Act, construction and regulation of buildings used for storage of foodstuffs must take into account public health concerns (sect. 101). The minister responsible may make rules for any purpose having as their object the preservation of health or the prevention of disease (sect. 104).

Uganda National Bureau of Standards Act, Chap 327

This act provides for the establishment of a national bureau of standards, the standardization of commodities and matters related thereto. A commodity is defined under the act to mean any article, product or thing which is or will ultimately be the subject of trade or use. The Uganda National Bureau of Standards (UNBS) is established as a body corporate under the general supervision of the minister responsible for commerce. The functions of UNBS include the formulation of national standard specifications for commodities; the promotion of standardization in commerce, industry, health, safety and social welfare; the endorsement or adoption of any
international or other country’s specification with or without modification as suitable for use in Uganda; the enforcement of standards in the protection of the public against harmful ingredients, dangerous components, shoddy material and poor performance; and the ability to seek membership of any international organization connected with standardization.

The governing body of UNBS is the National Standards Council (NSC), whose function is to declare standard specifications, certification marks and codes of practice and to do all things incidental thereto. The executive director of UNBS with the approval of the NSC may appoint standards inspectors to inspect and test any commodities and processes.

2.6. Biosafety

_Uganda National Council of Science and Technology Act (Chapter 209), 1990_

The Uganda National Council for Science and Technology (UNCST) is established as a body corporate under the general supervision of the minister responsible for planning and economic development. The functions of UNCST as stipulated in Section 4 of the act include:

- to advise on and coordinate the formulation of an explicit policy in all fields of science and technology;
- to assist in the promotion and development of indigenous science and technology through, _inter alia_, technology transfer and adaptation, as well as establishment of research and experimental development institutions, pilot plants and other testing grounds and standardization and quality control centres;
- to assist in the rationalization of the use of foreign science and technology;
- to act as a clearing house for information on research and experimental development taking place in scientific institutions, centres and other enterprises and on the potential application of their results;
- to work in close cooperation with and coordinate all scientific and technological activities of persons, institutions, sectors and organizations; and
- to carry out any other function incidental or conducive to these functions or as the minister may assign to it.

UNCST is authorized under Section 5 of the act to do all such things to facilitate its work or that are conducive or incidental to better carrying out its
functions. In this vein, UNCST may establish any specialized committees, research councils or organizations and carry out experimental and development activities or other scientific and technological services. UNCST may establish and maintain relationships with national, regional and international organizations and agencies as it may deem appropriate. There is established under the act a specialized committee on natural sciences whose mandate includes bio-science (sect. 15(1)(e)) and whose functions include advising UNCST on all policy matters on bio-science in the country; and advising on the assignment of scientific and technological responsibilities to different institutions or persons.

Represented on UNCST are the ministry responsible for agriculture, animal industry and fisheries; the ministry responsible for environment protection; the ministry responsible for health; UNBS; universities and eminent scientists in the field of agriculture and allied sciences, medical science and natural science.

UNCST established the National Biosafety Council (NBC) with members from the specialized departments/authorities of the various line ministries. The NBC is tasked with evaluating applications for confined field trials of living modified organisms (LMOs) and acting on referrals made by any department receiving applications for the import of LMOs (e.g. the Department of Crop Protection for seeds). While the decisions are made within the NBC, risk assessment is carried out by the competent departments/agencies.

UNCST, through its parent ministry, the Ministry of Finance and Economic Planning, has proposed the National Biotechnology and Biosafety Policy, June 2006. The policy defines the concepts of biotechnology and biosafety and the status of these concepts in Uganda. Biotechnology is defined as any technique that uses living organisms or substances therefrom to make or modify a product, improve plants or animals or microorganisms for specific purposes; and biosafety as the safe development, transfer and application of biotechnology and its products. The policy notes that the Uganda Biosafety Framework, 2000, by which various institutions have undertaken research in agricultural biotechnology and molecular biology, has limitations. The lack of an explicit biotechnology and biosafety policy has meant that national strategies and priorities in biotechnology development have not been proposed, and biotechnology considerations have not been integrated into the overall national development policy and planning framework.
The policy notes the inadequacy of the legal framework with respect to regulation of modern biotechnology and the issues that it raises. The legal provisions that exist are found in various pieces of sectoral legislation and are applied by a number of statutory bodies, each concerned with the fulfilment of its own mandate. Despite Uganda's ratification of the Convention on Biological Diversity in 1993 and the Cartagena Protocol in 2001, the provisions of these treaties have not been fully transformed into local laws nor is there an institution that can singularly address the concerns of these treaties.

The policy seeks to enable Uganda to realize the full potential of biotechnology through the formulation of a specific biotechnology and biosafety policy which defines the institutional, legal and regulatory regime for the promotion of biotechnology development and lays emphasis on infrastructure development, research, public awareness, human resource capacity development and the promotion of commercial/industrial development. The application of bioethics is required while government is encouraged to effectively integrate indigenous knowledge in the development and application of modern biotechnology. The policy proposes a monitoring and evaluation framework to continuously monitor and assess both the sector and system performance on the basis of measurable parameters.

III. INSTITUTIONAL BASIS FOR BIOSECURITY IN UGANDA: GAPS AND OVERLAPS

Biosecurity, as defined earlier in this report, is a strategic and integrated approach to analysing and managing risks in animal and plant life and health, food safety and biosafety. Issues arising within the Biosecurity framework are therefore cross-cutting and are the responsibility of different ministries, state agencies or departments.

The Ministry of Agriculture, Animal Industry and Fisheries (MAAIF) is responsible for animal health matters under its veterinary services division, while its fisheries arm handles aquatic life issues. Several inspectors of the veterinary services have recently received training on risk analysis but still face constraints in terms of equipment (e.g. laboratories).

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2 For a discussion of these instruments, see Chapter 2, Parts VI and VII.
The Department of Crop Protection is responsible for phytosanitary and plant protection matters and will be designated the national plant protection organization once the new law is enacted. The department has recently received technical assistance from FAO, which has resulted in the updating of the national pest list (still to be gazetted) and the effort to establish pest free areas for bananas.

The Ministry of Health (MOH) is the implementing authority of the Food and Drugs Act but it is commonly recognized that its human and financial resources are extremely limited and food safety is not prioritized. In practice, MOH has been working with other agencies on a case-by-case basis in instances where there have been emergencies (e.g. the European Union ban on fish from Uganda where MOH worked with MAAIF and UNBS to resolve the issues).

Inspection of food imports is mostly done by UNBS or its agents at entry points. Funding for food safety is sporadic although MOH seeks to have a systematic approach to food safety to address not only the final product stage but the production stage as well. Currently, MOH has a reactive approach to these matters. There is a very advanced food chain approach only for fisheries thanks to the cooperation between the Fisheries Department and UNBS.

Various other ministries are involved in Biosecurity matters mainly as supervisors of state agencies. The Ministry of Water and Environment houses the Wetlands Inspectorate which is the focal point for the implementation of the provisions of the Ramsar Convention. Agricultural research is undertaken by various agricultural research institutes that constitute the National Agricultural Research Organization, and by universities. Seed policy, certification standards and variety approval and release are the function of the National Seed Industry Authority.

NEMA as the principal agency responsible for the environment has a coordinating, monitoring and supervisory role. NEMA is the national focal point for the CBD and related instruments. The Uganda Wildlife Authority is the principal organ responsible for management of wildlife in Uganda. The National Forestry Authority has similar responsibility with respect to forests. UNBS, as the national standards body, sets and enforces standards, in some instances adopting standards from other jurisdictions or from international agencies for application in Uganda. The Uganda Revenue Authority, through
its customs department, plays a crucial role in ensuring the legitimacy of imports and exports of regulated materials. The Food Hygiene Advisory Committee and the responsible minister, in conjunction with the local governments and authorized persons, are responsible for the prevention of the adulteration of food.

UNCST is responsible for policy in all fields of science and technology. It is the competent authority for regulation and access to genetic resources, and is proposed in the draft policy on biotechnology and biosafety as the competent authority to supervise and regulate the implementation of the policy. Further, it is proposed as the competent authority for biosafety under the proposed Uganda Biosafety Bill, 2005, with the ministry responsible for the environment as the national focal point to provide coordinated communication on behalf of all relevant ministries, departments and agencies.

MAAIF through its crop protection and animal industry departments is responsible for the IPPC and OIE, respectively; and UNBS is responsible for the Codex Alimentarius. The Department of Crop Protection also acts as SPS Enquiry Point but has no visibility at the moment. It is advisable that this be addressed in the near future. NEMA and UNCST are the national focal point and competent authority for the CBD and Cartagena Protocol, respectively.

Each of the agencies mentioned above has inspectors charged with the duty of ensuring compliance with the provisions of each sectoral law. There are mechanisms in some laws authorizing delegation to related departments, thus enabling collaboration between departments and agencies. NEMA in gazetting environment inspectors is authorized to gazette persons employed as inspectors in other departments as environment inspectors, and such inspectors have been appointed. NEMA, however, lacks of human resources to participate in all activities of other ministries at the technical level.

Similarly, the phytosanitary service cooperates with the customs department in undertaking phytosanitary inspection at the various entry and exit points of Uganda whose borders are porous. Inspection of meat is undertaken by veterinary surgeons, medical officers or health inspectors authorized by the relevant minister or the local authority. However, most inspectors are only trained in the particular field of their employment and do not have sufficient capacity to effectively undertake inspection by delegation.

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3 For a discussion of the Codex Alimentarius, see Chapter 2, Part V.
Though UNBS is the national standards body, other government agencies or departments have the authority in law to set and enforce standards. NEMA sets environmental standards while the Directorate of Water Development sets water and water-related standards under the Water Act (Chapter 152, 1995).

**Biosecurity** as a concept has not been specifically addressed by policy or law in Uganda. NEMA as the principal agency responsible for the environment appears best placed to perform the umbrella role in biosecurity. NEMA already acts as a coordinating agency in various standing policy committees that are established (e.g. biodiversity, environment) and this focal point role is well accepted by other ministries. UNCST has the scientific and technical capacity to carry out the functions of a competent authority. Granted that UNCST at the moment is leaning towards biotechnology and biosafety, it could be persuaded to embrace **Biosecurity** as a whole.

The Plan for the Modernization of Agriculture (PMA) was approved by Cabinet in 2001, and includes the strengthening of plant and animal health controls. Nine ministries are directly involved with PMA whose main function is the commercialization of agriculture in order to eradicate poverty in Uganda. The PMA process of mainstreaming issues involves a Stakeholders Forum, a Steering Committee comprised of Permanent Secretaries and the PMA Secretariat. The PMA operates through consensus building through annual reviews of the PMA. All the responsibilities for implementation of the PMA remain with the ministries. The Stakeholders Forum and the Steering Committee seem to be the appropriate bodies where stakeholders and policy makers can be sensitized to **Biosecurity**.

It is important to create public awareness about **Biosecurity** and what the cost of a disease outbreak would be to the country. This would engender better understanding and policing of already existing laws, and improve the efficiency of mandated institutions even more.

**IV. CONCLUSIONS AND WAY FORWARD**

The National Biotechnology and Biosafety Policy is in the final stages of consideration in the Ministry of Finance. The institutional framework proposed in the policy reflects some of the institutional overlaps that exist in the current operations as pointed out in the review of the national legal regime. A National Biosafety Act is proposed but is still in the initial stages of formulation. The underpinning philosophy of the act is that biosafety is a
cross-sectoral activity spanning from food safety, plant and animal health to environmental protection and thus requiring operative coordination among the authorities. Biosecurity has the same characteristics. Uganda has the opportunity to incorporate in policy and law an appropriate coordination mechanism for Biosecurity.

Biosecurity is not the responsibility of one agency of state; as seen above, it involves several ministries, departments and agencies. Accordingly the regulatory framework for Biosecurity must be multi-sectoral in nature with an overall coordinating body. Any other formulation would engender institutional rivalry and conflict leading to paralysis.

Several pieces of legislation which address food safety, plant and animal health have been discussed. The legislation is sectoral in nature particularly since different departments or regulatory agencies are responsible for the implementation of each law. The Plant Protection Act has been reviewed to bring it in conformity with the IPPC and may be further reviewed to ensure compliance with new developments in Biosecurity. The National Environment Act as the framework law on the environment does concern itself with CBD matters though there are sectoral laws in force that address CBD matters in greater detail, for instance the Uganda Wildlife Act. The proposed National Biosafety Act which seeks to domesticate the Cartagena Protocol is being formulated under the ministry responsible for science and technology which at the moment is the Ministry of Finance. This in itself poses a challenge since the Ministry of Finance has as its priority the fiscal matters of state and macro- and other economic issues. The elaboration of a new Food Act is at a very early stage. All these efforts show a certain degree of commitment to aligning the national legal framework to international dictates. In that respect, Biosecurity may become the guiding principle for legislative upgrading and give new impetus to the various legislative initiatives.

In the development of an appropriate legal and institutional framework for Biosecurity, it is recommended that the National Biosafety Act serve as a means of ensuring coordination and cross-sectoral management in Biosecurity. The government policy does not encourage the establishment of new institutions at the moment. In the circumstances, it is better to encourage the various institutions to carry out their mandates with Biosecurity in mind. In that regard, the cross-sectoral coordination that is being promoted for biosafety represents the ideal occasion to mandate the various institutions to embrace Biosecurity as a whole.
VIET NAM COUNTRY STUDY*

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* This chapter was prepared by Duong Thanh An.
I. INTRODUCTION

FAO uses the term Biosecurity in relation to sanitary, phytosanitary and zoosanitary measures applied in food and agriculture regulatory systems. For FAO, Biosecurity broadly describes the process and objective of managing biological risks associated with food and agriculture in a holistic manner. Biosecurity is a holistic concept of direct relevance to the sustainability of agriculture, food safety and the protection of the environment, including biodiversity. This chapter reviews Viet Nam’s legislation in the five areas of Biosecurity, namely, food safety, plant health, animal health, IAS and biosafety.

II. VIET NAM’S LEGISLATION ON BIOSECURITY

2.1. Food safety

In Viet Nam, food safety is regulated mostly by the 2003 Ordinance on Food Hygiene and Safety and other related laws, namely, the 2000 Law on People’s Health, the 2003 Fisheries Law and the 1999 Ordinance on Consumer Protection.

Ordinance on Hygiene and Food Safety (2003)

The Ordinance on Food Hygiene and Safety (OFHS) establishes a regulatory regime for food safety in Viet Nam. It includes provisions to ensure hygiene and food safety in food production and trade as well as prevention and control of food poisoning and food-borne diseases.

In the area of food production and trade

According to the OFHS, production and trade in food include farming, harvesting, treating, processing, packaging, keeping in storage and transporting. The ordinance provides for: (1) standards for hygiene and food safety; (2) regulations on conditions of food production and trade; (3) regulations on food processing; (4) regulations on storage and transport of food; and (5) regulations on export and import of food.

The Vietnamese standards on hygiene and food safety include national standards, ministerial standards and local standards. National standards are established by the Ministry of Science and Technology (MOST) in cooperation with the Ministry of Health (MOH) and other related ministries.
Ministerial standards are established by other ministries, such as the Ministry of Agriculture and Rural Development (MOARD), while local standards are developed by authorized food production enterprises in accordance with national and ministerial standards. Organizations and individuals producing and trading in food are required to publish the standards that they follow.

The OFHS provides that organizations and individuals shall comply with three sets of regulations:

- safety regulations on infrastructure, such as locations, water supply systems and waste water treatment;
- regulations on equipment, such as processing, storage and transportation facilities; and
- regulations on personnel, such as employees’ health and knowledge of hygiene and food safety principles.

Enterprises that produce and trade in fresh food are required to ensure that the production and trading places are clean and isolated from polluted areas. They are also responsible for applying appropriate methods of storage and transport.

Regulations on food export and import require enterprises to obtain an authorization from public authorities certifying that they have adequate food safety management infrastructure. Where the requirements are not met, food may be seized and disposed of.

Decree No. 163/2004/ND-CP of the Government of Viet Nam dated 7 September 2004 provides for the implementation of the OFHS. The responsibilities of various ministries are specified in articles 21–29 as follows:

- MOH is responsible for formulating and promulgating strategies and policies on food hygiene and safety. Also, MOH must assume the lead in coordinating with other ministries and ministerial departments in: (1) implementing food safety controls in the retail sector and testing pesticide residues in food; (2) inspecting and testing on food hygiene; and (3) carrying out research, training, international cooperation and awareness raising.
- MOARD is responsible for the production process, including processing, slaughtering, preservation and transport as well as for veterinary controls on imported food of animal origin.
• The Ministry of Fisheries (MOF) is responsible for aquatic products for domestic consumption and aquatic products which are exported or temporarily imported for re-export.
• The Ministry of Industry (MOI) is responsible for food products originating from establishments under its management.
• The Ministry of Trade is responsible for food safety controls in the retail sector and, in coordination with other ministries and ministerial departments, for the regulation of food businesses.
• MOST is responsible for standard setting, quality controls and licences, in coordination with MOH.
• The Ministry of Culture and Information is responsible for the regulation of food advertising and for awareness raising on food safety, in coordination with MOH.
• The Ministry of Finance is responsible for the collection of fees and charges and the inspection of food for import in collaboration with MOH and in accordance with customs legislation.
• The People’s Committees at the different territorial levels are required to assist state authorities with the implementation of food safety and are responsible for implementing good manufacturing practices and building models of community participation in the management of food hygiene and safety.

In addition, all the ministries shall formulate and promulgate regulations in their respective areas of competence in coordination with MOH.

In the area of prevention and control of food poisoning and food-borne diseases

When food poisoning occurs or food-borne diseases occur at a specific location, the local People’s Committee is responsible for applying measures to prevent the transmission of disease and informing the public. In cases where the outbreak is in a large area and seriously threatens public health, emergency regulations must be taken into account.

Additional responsibilities to be assumed by the relevant ministries are as follows:

• as for prevention, MOARD and MOF shall implement good manufacturing practices in order to ensure hygiene and safety for
agricultural and aquatic products before they are marketed, while MOI has the same responsibilities for food production sites;

- MOH is responsible for the implementation and enforcement of food safety and hygiene standards as well as the management of food safety emergencies.

Food producing and food trading organizations as well as individuals shall observe food safety and hygiene regulations and standards. In cases of food poisoning or food-borne diseases, they must immediately report to the local health administrations and take remedial measures as instructed. For penalties, depending on the seriousness of violations, individuals and legal entities shall be sanctioned administratively or through penal liability and shall pay compensation.

*Law on the Protection of People’s Health (2000)*

Although the Law on the Protection of People’s Health mostly regulates issues of public health protection, it also deals with some aspects related to food hygiene and safety. First, the law states that Vietnamese citizens have the right to health protection as well as the right of access to safe and wholesome food.

Second, article 7 provides for regulations on hygiene of food, water and alcohol. Enterprises that produce, process, store or transport these items must follow certain hygiene standards. The use of chemicals for food processing and preservation without the permission of MOH is prohibited. Individuals who have transmissible diseases are banned from activities in direct contact with food.

Third, hygiene in animal farms, an important factor of food safety, is also covered by article 11 of this law. It prohibits slaughtering, trading in and consuming livestock or poultry that may bear transmissible diseases.

*Fisheries Law (2003)*

The Fisheries Law has some regulations related to food safety. Chapter VI regulates processing activities of aquatic products for which the following two requirements are set out:
• processing enterprises of aquatic products are required to have their own storage areas as well as processing and hygiene equipment which shall meet certain technical and hygiene standards. Also, the use of additives and chemicals that are in a banned list is prohibited; and
• enterprises that process, import or export aquatic products are responsible for ensuring conformity with processing and product standards.

MOF, in coordination with other relevant ministries, is responsible for inspecting facilities and monitoring compliance with regulations relating to quality and safety of imported, exported or domestically produced aquatic products.

Ordinance on Veterinary Controls (2004)

The Ordinance on Veterinary Controls provides for standards on slaughter to ensure the safety of food of animal origin. Article 8 provides that all activities of slaughtering in slaughterhouses which do not meet certain hygiene standards are prohibited. Also, the slaughtering of infected animals and the trading in infected food of animal origin are prohibited.

With regard to oversight of slaughter, the ordinance has three main prescriptions:

• all slaughtered animals must be inspected to make sure the standards on hygiene and food safety are met;
• slaughterhouses and processing places must meet standards of hygiene and safety;
• slaughterers must be in good health and bear no transmissible diseases as well as undergo periodic health check-ups by local public health authorities.

Article 7 of the ordinance provides that veterinary standards include Vietnamese standards, professional standards, corporate standards and international standards applicable to Viet Nam. MOST is responsible for promulgating Vietnamese standards while MOARD and MOF promulgate professional standards. Corporate standards are developed by private establishments operating in the sector.
Ordinance on Consumer Protection (1999)

Among the objectives of the Ordinance on Consumer Protection are the protection of the health of the general public. It reaffirms the duty of individuals and enterprises to follow hygiene standards and establishes administrative and criminal offences and penalties. Pursuant to the ordinance, MOH, MOF and MOARD are the implementing authorities.

Other instruments

Decree No. 21/2006/ND-CP of the Government of Viet Nam dated 27 February 2006 promulgated regulations on the trade in and use of nutrition products for children. The objective of the decree is to ensure that all children are protected from unsafe nutrition products. MOH is required to cooperate with other relevant ministries to promote the use of breast milk. All nutrition products for children must comply with standards developed in accordance with food safety laws.

Decision No. 43/2006/QD-TTg of the Prime Minister dated 20 February 2006 approved the National Action Plan on Hygiene and Food Safety to 2010. The plan aims to ensure food safety in order to protect human health and facilitate socio-economic development. Eighty percent of national standards are expected to be in conformity with international standards by 2010.

Decision No. 21/2007/QD-BYT of MOH dated 12 March 2007 promulgated regulations on the health of operators directly in contact with food during the processing of pre-packaged food and trading in instant food.

Assessment of the legislation

The above analysis shows a relatively complete regulatory framework covering the main areas of food safety. In particular, it is worth noting that the government is committed to progressively achieving compliance with international standards in order to protect human health and promote trade.

However, it must be noted that, although laws are in place, the regulatory activities have limited impact on the general food safety situation in the country and the implementation arrangements are not efficient. This might be caused by the following factors:
First, general laws are in place but the implementing regulations are inadequate or unapplied. Below are some illustrative examples:

- manufacturers of and traders in meat and meat products, eggs and egg products and other food products are obliged to obtain a licence and a certificate from MOH. However, street food vendors are not subject to the regime. Moreover, MOH has authorized local medical centres to issue health and professional certificates to food dealers. However, many of those centres lack capacity to implement this function effectively;
- the state authority responsible for the management and inspection of food safety is MOH, while the state authority responsible for performing quality controls is MOST. This division of responsibilities has led to undesirable results and harmed efficient food safety management as the two ministries do not coordinate on a regular basis;
- the maximum administrative penalty of 15 million dong for the production of and trading in unsafe or poisoned food is low (approximately €660 at current rates), given the financial capability of major food enterprises;
- the national food safety standards are permissive and many low-quality food items have entered the market with negative and uncontrolled consequences on food safety; and
- under the Ordinance on Consumer Protection, consumers who purchase products not conforming to regulations and standards can only file a complaint with the Association of Consumer Advocates, which is a non-state entity without any power to adjudicate disputes or order compensation.

Second, the coordination of activities among government authorities on food safety is still weak. Since food comes from multiple origins, food safety is related to the management functions of several governmental sectors such as public health, veterinary services and fisheries. However, the clarification of their precise responsibilities along the food chain, from primary production to consumption, has not been achieved yet. The laws contain general provisions that are subject to interpretation. As a result, in the implementation phase the activities of the sectoral authorities overlap. In some cases, those authorities claim a more extended mandate for food safety than is established in the relevant legislation. In some other cases
(e.g. food emergencies), there are gaps in regulatory activities. Besides, many authorities have limited enforcement and infrastructure capabilities.

Finally, there is not enough information on food safety channelled to the public, hence citizens lack necessary information to become "knowledgeable consumers".

2.2. Animal health

Legal provisions on the protection of animal health and life are found in different laws, as follows:


The Fisheries Law regulates activities related to aquatic animals and aquatic animal products, such as breeding, processing, import and export. Activities that cause adverse effects on aquatic animal breeds are generally prohibited. The law establishes a list of aquatic animal species for which aquaculture is prohibited as well as a list of chemicals that are banned in aquaculture. The law envisages a series of measures to protect the living environment for aquatic animals as well as to preserve endangered species. Aquaculture of endangered species requires the permission of MOF or the provincial People’s Committee.

Individuals who breed aquatic animal species shall comply with regulations on breeding and the use of chemicals. The law also speaks of animal health measures to prevent the outbreak of or to control the spread of animal diseases in aquaculture environments.

Article 35 of the law states that MOF is responsible for developing: (1) standards for feed used in aquaculture; (2) zoosanitary measures in aquaculture; and (3) the list of banned chemicals. The responsibilities for prevention and control of animal diseases in aquaculture environments are assigned to MOF and the provincial People’s Committees.

*Ordinance on Veterinary Controls* (2004)

Article 9 of the Ordinance on Veterinary Controls provides for:

- surveillance and control;
• animal quarantine and zoosanitary inspections;
• programmes on control and eradication of animal diseases and zoonoses; and
• quality control of products of animal origin, animal feed, veterinary drugs, veterinary biological products and microorganisms.

All activities that cause adverse effects on animal health are prohibited, including non-compliance with regulations on disease prevention, disposal of carcasses and movement of infected animals and animal products. MOARD and MOF are responsible for the prevention and control of animal diseases, including the treatment of infected animals. The government is mandated to establish a National Steering Committee for animal disease prevention and control, upon request of MOARD or MOF.

MOARD, MOF, the People's Committees at different territorial levels, government veterinarians, customs officials and officials of transport authorities are collectively assigned the responsibility for animal quarantine. Animal quarantine includes the quarantine of domestic animals and animal products, animals and animal products for import and export as well as animals and animal products in transit. Article 23 of the ordinance provides that all animals and animal products must, when being transported out of their district of origin, be quarantined at departure. Also, articles 28 and 29 of the ordinance have regulations on quarantine for imported and exported animals and animal products. Article 26 establishes requirements for animals and animal products for domestic transportation.

The ordinance gives the responsibility for the management of veterinary drugs and veterinary biological products, including microorganisms, to MOARD, MOF and the People's Committees.

Ordinance on Livestock Breeds (2004)

The Ordinance on Livestock Breeds has some provisions related to animal health and life. It generally prohibits activities that may harm safe animal breeding and regulates some zoosanitary aspects of animal breeding and multiplication. Article 9 of the ordinance prohibits the export of livestock species of genetic value. MOARD is responsible for the state management of agricultural livestock breeds while MOF is responsible for aquatic livestock breeds.
Assessment of the legislation

As presented above, the regulatory framework on animal health and life assigns responsibilities to authorities and individuals in general terms, for instance by assigning the mandate for certain activities to ministries or by setting forth a general prohibition of harmful conduct. In cases where individuals do not comply, administrative and criminal offences and penalties are in place. The criminal law of Viet Nam (1999) contains applicable provisions, such as those in articles 158 and 187.

Certain shortcomings of the legislation can be identified:

- the legal dictates are of a very general nature and some powers of public authorities (e.g. inspections by veterinarians) and duties of individuals (e.g. duties of animal owners) are not legislated;
- small animal husbandry is very relevant in Viet Nam but some key regulatory activities, such as inspection of cattle and slaughter, do not reach small household farms;
- the task force of veterinarians and animal health inspectors within MOARD is still deficient, with limited professional knowledge and infrastructure; and
- information exchange between MOARD and the local People’s Committees is not efficient and, in cases of epidemics, this causes delays in control measures.

2.3. Plant health

In addition to the phytosanitary legislation listed in Chapter 3, two other laws, not specific to plant health, contain scattered provisions referring to plant quarantine. The Law on Environment Protection (2005) generally prohibits the import of plants without an import permit, the issuance of which is administered by plant health legislation. The 2004 Law on Forest Protection and Development refers to regulations on the prevention and eradication of plant pests and requires individuals to execute control measures in accordance with the guidelines of state authorities. The law also refers to import and export requirements that are established in plant protection legislation.

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1 This part of the report draws from FAO, Technical Assistance in Phytosanitary Legislation, Vietnam, 2005.
2 See Chapter 3, Section 3.6.
Institutional mandate of the Plant Protection Department

The Plant Protection Department (PPD) was established in 1961 as the authority responsible for plant protection and plant quarantine matters in Viet Nam. It is affiliated with MOARD. The actual mandate of the PPD is set forth in Decision No. 88/2003 of MOARD.

The three main areas of the PPD’s mandate are: (1) plant quarantine; (2) plant protection; and (3) pesticides management. With regard to plant quarantine, article 5(b) of Decision No. 88 speaks of: (a) administration of plant quarantine activities; (b) elaboration of the list of quarantine pests for approval by the minister responsible for agriculture; (c) control of pest outbreaks; (d) phytosanitary inspection for imports and exports; and (e) treatment of consignments.

Under article 5(a), PPD exercises the following responsibilities in connection with plant protection: (a) surveillance for the purpose of reporting the occurrence of pests; and (b) proposals to the minister for the declaration of quarantine areas.

General responsibilities affecting phytosanitary activities include scientific research; policy-making for trade in plants and plant products; international cooperation and representation of the country in international fora; implementation of relevant international agreements; and training of staff.

The mandate of "Plant Quarantine Agencies" is set forth in article 5 of Decree No. 02/2007/ND-CP on Plant Quarantine. Those provisions make an important addition to the PPD mandate, namely, the designation and management of pest free areas.

Surveillance and pest control

Article 9 of the Ordinance on Plant Protection and Quarantine (2001) generally refers to the management of injurious pests, including survey, detection, forecasting and warning of pest occurrence, development, distribution and damage.

Article 10 specifies the rights and duties of plant resource owners which include: (1) the right to be informed on pest status and assisted with pest control by the competent governmental bodies; (2) the duty to apply
appropriate pest control measures as recommended by competent governmental bodies in order to contain a pest; and (3) the duty to report any pest of economic importance to competent governmental bodies.

Declaration and management of quarantine areas

Article 11(2) of the ordinance sets forth the mandate for designation of areas where an outbreak of a pest of economic importance occurs. The mandate lies with the Chair of the People’s Committee or MOARD depending on the territorial extension of the outbreak. As for management of affected areas, article 12(1) tasks MOARD with ordering and implementing control measures in collaboration with local authorities. Provisions made in article 16 of the ordinance state the duties of individuals to report and cooperate in the implementation of control measures in cases of pest outbreaks.

With regard to emergency action, article 11(1) of the ordinance provides that, when a pest outbreak is reported, the relevant organizations shall inspect promptly and assist the owner of the plant resources with pest control. Article 17 states that when quarantine or alien pests are detected, competent plant health authorities shall decide upon appropriate measures to delimit and eradicate such pests and request the owners of regulated articles to undertake those measures immediately. Those provisions appear open to interpretation in cases of emergency (i.e. in situations where prompt phytosanitary action is undertaken in a new or unexpected phytosanitary situation without full technical justification).

Import controls

Article 14 of the ordinance generally describes phytosanitary activities related to imports, which include inspection, detection, treatment and monitoring of the status of the consignment after import. Article 18(1) prescribes that all regulated articles for import are subject to phytosanitary inspection while article 22 sets forth a corresponding duty for importers and establishes that, in cases where a quarantine pest is detected, the plant health authority can order re-export, destruction, observation or treatment of the consignment. In terms of article 13(d) of Decree No. 02 on Plant Quarantine, MOARD is responsible for issuing permits for the import of certain beneficial organisms to be specified in a list.
Article 19 of the ordinance sets out criteria for the import of seeds as follows: (1) seed imports are subject to strict inspection and monitoring by the plant health authority; (b) movement of consignments shall be traceable; (c) seeds that are imported for the first time shall be grown in a designated place for phytosanitary testing and shall be released only after certification of freedom from quarantine pests.

**List of regulated pests and articles**

Article 15 of the ordinance and article 7(1) of the decree provide that MOARD shall publish and regularly update the list of quarantine pests and the list of regulated articles. Decision No. 88 tasks the PPD with elaboration of these lists.

Decision No. 117/2000 of MOARD contains the list of quarantine pests of Viet Nam, which are divided into two categories: (a) pests of potential economic importance and not present in Viet Nam; and (b) pests of potential economic importance which are present in the territory but not widely distributed. These definitions mainly track the terminology and concepts of the new revised text of the International Plant Protection Convention.3

Decision No. 56/2001 of MOARD provides the list of regulated articles for import, export, re-import and re-export including plant seeds, plants and parts thereof, plant products, insects, diseases, weeds, soil and other materials harbouring pests, and means of conveyance.

**Phytosanitary certification for export**

Article 20 of the ordinance and article 15 of the decree indicate that phytosanitary inspection for export and re-export shall be carried out on regulated articles in cases where it is so required by commercial contracts or international agreements to which Viet Nam adheres or where so requested by the exporter. In cases of non-compliance with the phytosanitary requirements of the importing country, phytosanitary certification shall be refused until the exporter properly treats the consignment.

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3 For a discussion of this instrument, see Chapter 2, Part III.
Article 16 of the decree specifies the procedures for export inspections, while article 17 mandates the plant health authority to monitor the status of consignments after certification and prior to export.

With regard to consignments in transit, article 21 of the ordinance prescribes mandatory phytosanitary inspection and action in cases of infestation while articles 18 and 19 of the decree detail procedures for inspection of consignments in transit.

Miscellaneous

With regard to powers of quarantine officers, article 6(3) of the decree empowers quarantine officers to enter any place where regulated articles are found. Offences and penalties are regulated in subsidiary legislation, namely, Decree No. 26/2003/ND-CP.

Assessment of the legislation

The legislation in place comprehensively covers most of the operational areas of the PPD at present. Its mandate, as set out in Decision No. 88, is well reflected in the substantive provisions of the ordinance. The ordinance also has a prescriptive force in that it imposes duties on individuals (e.g. duty to notify the presence of a pest, duty to apply for permits and certificates, duty to pay fees).

However, the functioning of the PPD may suffer from a number of shortcomings arising from the legislative framework. At first glance, the absence of parliamentary-level legislation is striking, in terms of protection of rights as well as definition of institutional and individual responsibilities. A well-grounded legislative framework establishing predictable rules for phytosanitary controls cannot be established without parliamentary-level legislation.

The provisions for administration of the ordinance and the decree do not directly task the PPD with any responsibility. This may not be efficient in light of the designation of the PPD as the national plant protection organization for Viet Nam, which is a requirement of the IPPC.

At the level of substantive provisions, the legislation provides for some key principles (e.g. pest risk analysis) and regulatory areas (e.g. pest free areas)
which are dealt with in the international agreements. With the recent adoption of the decree, controls on imports and exports, which are important regulatory areas from the perspective of international trade, are disciplined with clarity.

2.4. Invasive alien species

Invasive alien species (IAS) are not systematically addressed in Viet Nam’s legislation. Some IAS have been imported for commercial purposes but it is only in relation to some of them that the National Environment Agency has conducted an assessment of their impacts on biodiversity and the environment. The concept of IAS also appears sporadically in regulations on biodiversity, plant and animal health.

Decree No. 58/2002/ND/CP of the Government of Viet Nam dated 3 June 2002 includes provisions on plant quarantine. Article 16 establishes that the import of all IAS of plant origin is prohibited. In specific cases where the import is for scientific purposes, the permission from the Minister of Agriculture and Rural Development may be sought.

Decree No. 109/2003/ND/CP of the Government of Viet Nam dated 23 September 2003 contains provisions on the preservation and sustainable development of wetlands. In this decree, the introduction of new species which may endanger the ecosystem or alter the gene pool of animals and plants in these areas is banned.

Article 6 of the Fisheries Law (2003) states that the farming of new aquatic animal species without permission from MOF is prohibited. Individuals and organizations may farm aquatic animal species that appear in a list of permitted species.

The Ordinance on Plant Breeding (2004) prohibits the import, breeding and commercialization of IAS that may cause harm to human health, environment and ecosystems. The Ordinance on Livestock Breeds contains similar provisions.

The Ordinance on Veterinary Controls (2004) provides that all animal species that cause harm to human health, animal, environment, and ecological system are subject to quarantine.
The Law on Forest Protection and Development (2004) establishes a permit system for exploitation activities in protected forestry areas. The permit system could operate to manage IAS in those areas.

Thus, the laws of Viet Nam sporadically address IAS in different instruments. Accordingly, the institutional responsibilities are assigned to different state authorities without any overarching authority. It is critical to develop a new framework to manage and effectively control IAS under a designated authority responsible for their management. Specifically, this institution should be responsible for investigating, categorizing and conducting surveillance of IAS. The authority should also be tasked with the development of risk assessment procedures to evaluate applications for the import of IAS.

2.5. Biosafety

Until 2004, the issue of management of genetically modified organisms (GMOs) was not properly addressed in Viet Nam’s legislation. It was sporadically touched upon in some laws, such as Decree No. 109 (referred to in the preceding section), the Ordinance on Animal Breeding, the Ordinance on Plant Breeding, Decision No. 178/1999/QD-TTg on the Labelling of Domestic and Import-export Goods and the Ordinance on Hygiene and Food Safety. Most of those laws generally mentioned GMOs simply to include them within the scope of the legislative instrument, but contained no specific directives on this issue.

Regulation of biosafety in environmental legislation

The Law on Environment Protection (2005) includes several provisions on biosafety. The purpose of those provisions is to extend and apply the existing legislative framework for conventional processes and products to biotechnology, GMOs and GMO products. Article 87 provides as follows:

- individuals and legal entities that manufacture and trade in GMOs and their products shall comply with the laws on biodiversity, food hygiene and food safety, plant and animal breeding and other related laws;
- Individuals and legal entities are only permitted to carry out research, manufacture, trade in, use, import, export, storage and transportation of GMOs and their products which are included in a list. Those activities shall be in compliance with legislation in place; and
• genetically modified animals, plants and microorganisms for import shall be quarantined.

Regulations on the management of biosafety were promulgated by Decision No. 212/2005/QD-TTg of 26 August 2005. The regulations provide for several areas of biosafety management, from research, manufacturing and trade, to importation, exportation, storage and transportation of GMOs.

The Ministry of Natural Resources and Environment (MONRE) is the principal authority responsible for management of biosafety at the state level. Other ministries, such as MOF, MOARD and MOH, are responsible at the ministerial level. According to regulations issued in 2005, enterprises shall obtain a biosafety certification for their risk management measures. For all other aspects of biosafety (e.g. authorization to import, risk assessment), the regulations lack detailed provisions.

MONRE has a coordinating role vis-à-vis the other ministries. With regard to conservation and biodiversity, Decree No. 109 regulates the prevention of adverse effects of GMOs on wetland ecosystems. Article 7 prohibits "the introduction of new species into wetland ecosystems which may cause ecological unbalance or modify the gene pool of local animals and plants". In the implementing Circular No. 18/2004/TT-BTNMT, there is no specific directive on how to manage biosafety in those ecosystems.

It is clear from the above that biosafety has been integrated into the Law on Environment Protection and other subsidiary legislation in the area of environment law, but in very general terms without any operational detail or implementation arrangements.

Regulation of biosafety in plant and animal legislation

The plant and animal legislation of Viet Nam follows the same approach as the environmental laws, which means applying legislation developed for conventional agriculture to biotechnology. The Ordinance on Animal Breeding and the Ordinance on Plant Breeding state that activities of research, selection, manufacturing, trading in, import and export of genetically modified animal and plant varieties shall be in compliance with applicable laws.
The 2001 Ordinance on Plant Protection and Quarantine does not directly regulate GMOs. However, pest risk analysis carried out by the PPD under the ordinance and in accordance with international standards may cover GMO plants and plant products. All other regulatory activities of the PPD could apply to genetically modified plants and plant products as well.

Regulation of biosafety in food safety legislation

Under the Ordinance on Hygiene and Food Safety, the issue of products and food originating from GMOs is sporadically covered. Article 20 provides that products and food originating from GMOs must clearly be labelled as such in Vietnamese. Decision No. 178 on Labelling (referred to above) provides some directives to implement the ordinance.

Ministries and their departments have promulgated other legal directives on labelling. MOF and MOARD promulgated Circular No. 03/2000/TT-BTS on 22 September 2000 and Circular No. 102/2001/TT-BNN-KHCN on 26 October 2001, respectively. These directives provide that food originating from GMOs must be labelled clearly in Vietnamese.

Currently, a draft Law on Biodiversity is under development. It covers biosafety in that it provides that MONRE is the designated implementing authority for the Cartagena Protocol.4

III. ASSESSMENT OF LEGISLATION

It is clear from the above that Viet Nam does not have a single law which covers the whole Biosecurity issue. Instead, provisions on Biosecurity are found in different laws and regulations which address specific areas of Biosecurity. The institutional mandates of the ministries that play a role in the Biosecurity scenario are briefly summarized below.

MOARD plays the most important role in the area of animal and plant health. It establishes and implements quarantine measures for animals and plants. In the food safety area, it is responsible for the production processes of food and the management of hygiene of imported food of animal origin. In addition, it serves as enquiry point and notification

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4 For a discussion of this instrument see Chapter 2, Part VII.
authority under the World Trade Organization (WTO) Agreement on the Application of Sanitary and Phytosanitary (SPS) Measures.5

MOF is responsible for sanitary measures for aquatic animals and animal products. In the food safety area, it is responsible for aquatic animal products for domestic consumption and aquatic animal products which are exported or temporarily imported for re-export. MOF, in coordination with other relevant ministries, is responsible for inspecting enterprises and monitoring compliance with regulations on the quality and safety of imported and exported aquatic animal products. According to a recent resolution of the National Assembly, from August 2007 MOF will be merged with MOARD.

MOST is responsible for promulgating standards in all areas of Biosecurity. It is also the Codex Contact Point.6

MONRE is responsible for the management of biosafety at the overarching state level. The Environment Protection Agency that operates under its hierarchy is the focal point for the Convention on Biological Diversity7 and the Cartagena Protocol.

MOH is responsible for formulating and promulgating strategies and policies on food hygiene and safety as well as taking an oversight role in the prevention of food poisoning. It is in charge of the development of regulations on food hygiene and safety and their enforcement. In cases of emergencies, it coordinates with the People’s Committees at the different territorial levels as well as the concerned ministries to establish control measures.

In short, in the five areas of Biosecurity, the existing laws of Viet Nam have tasked several ministries with the performance of functions. In terms of substantive regulation of Biosecurity, the plant health framework seems to be the most advanced. The legislation captures the principles and the key regulatory areas according to international dictates. In light of the WTO SPS requirements by which Viet Nam has recently become bound, it is critical to have enabling laws and regulations in the other areas of Biosecurity. Biosecurity is the guiding concept that can bring together the existing and future legislative efforts in order to manage trade in agricultural products in a manner respectful of the SPS dictates while at the same time protective of the natural resources of the country.

5 For a discussion of this agreement, see Chapter 2, Section 2.1.
6 For a discussion of the Codex Alimentarius, see Chapter 2, Part V.
7 For a discussion of this convention, see Chapter 2, Part VI.
CONCLUSION

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I. LESSONS LEARNED

*Biosecurity* is an evolving concept not only because of progress in science and technology that brings new opportunities and new risks to food and agriculture but also because of how the approach is perceived and implemented in different countries. Governments give varying degrees of priority to international trade and to the protection of agricultural resources from sanitary and phytosanitary threats, thus *biosecurity* is addressed differently in different jurisdictions. In some countries, *biosecurity* functions as a specific objective of agricultural policies and laws, and governments take specific action to synchronize sectoral authorities and responsibilities; in other countries interest in a *biosecurity* approach can simply point to the need to protect agricultural health and food safety against trade liberalization without requiring any immediate action. What is agreed is that the path to a shared perception of *biosecurity* and its legal framework is still long.

On the legislative side, *biosecurity* does not appear as an integrated whole in any of the pilot countries. Rather, *biosecurity* is addressed through sectoral regulatory instruments that may have arisen in different historical contexts and because of specific needs. In Chapters 5–10, the national legal consultants examined these legislative instruments in an attempt to evaluate their compliance with international norms and their effectiveness at coordinating action against biological risks. The consultants essentially pieced together the sectoral regulatory instruments of different scopes and examined them from a new angle, assessing how well these pieces of legislation embrace *biosecurity*. Their work reveals gaps in certain sectors as well as the absence of a consistent approach to *biosecurity* overall.

At the institutional level, the national studies show that coordination is often lacking among government bodies involved in *biosecurity* matters, and inter-institutional conflicts are common. This is an issue in the six countries reviewed and argues for some corrective action. If the main goal of *biosecurity* at national level is to integrate the animal health, plant health and food safety sectors, the national studies reveal that this dimension of *biosecurity* is not being successfully achieved.

Regardless of whether the governments of the pilot countries have overtly committed themselves to the implementation of a *biosecurity* approach, the case studies reveal that all six countries are making efforts to align their legislation with international dictates in some or all of the areas that
constitute Biosecurity. In that respect, Biosecurity has in some fashion already become the guiding principle for legislative upgrading. Greater understanding can provide further impetus and coherence to many legislative initiatives already under way.

At the international level, Biosecurity is not defined in any single legal text, instead covering a range of subjects and implicating several international instruments. Governments may wish to consider all these instruments under the Biosecurity rubric in order to implement their international obligations in a coordinated manner. This should help countries protect their natural resources for food and agriculture more effectively.

II. THE WAY FORWARD

The rapid growth of international trade in agricultural products calls for prompt action at the national level to avoid biological risks. As can be seen from the preceding chapters, any legislative and institutional reform must be supported at the political level. This is particularly true in a cross-cutting area like Biosecurity which requires a coherent approach.

In some countries, even where a comprehensive policy and clear strategy may be adopted, legal reforms may be blocked for years at the final drafting stage before draft legislative instruments are submitted to parliament (for primary legislation) or to the relevant minister or ministers (for subsidiary legislation). Where no political reasons are hindering adoption of the text, there may be staffing problems among the legal personnel in charge either of drafting the necessary legislation or of checking its consistency with the domestic legal system.

This problem has been noted in Kenya, where the Law Reform Commission has been trying to consolidate the legislative framework for agriculture, which consists of more than 130 legal instruments. The consolidation of legislation (i.e. the reduction in the number of legislative enactments by merging several instruments dealing with common issues into one piece of legislation) is seen as a way of fast-tracking and updating laws in a context where legislation takes a long time to pass parliament. This is also a perfect opportunity to review the laws through the lens of Biosecurity.

Another important task at national level will be to evaluate the actual capacity of the institutions that will be called upon to enforce Biosecurity
legislation. Without such an assessment, the application of the legal methodology contained in this study and the development of a strategy for legislative and institutional reform risks becoming a theoretical exercise. Capacity-building and provision of essential resources, based on a real assessment of existing institutions and capacities, are likely to be essential to the implementation of an effective Biosecurity approach.

On a more practical note, two final observations can be made. First, it appeared from the national consultations and from the authors’ extensive experience in FAO member countries that Biosecurity is not a particularly popular or well-understood concept as yet. It is often confused with biosafety, which has been the subject of an international convention as well as capacity-building initiatives worldwide. This suggests that an important next step will be to carry out awareness-raising activities at national and regional level. Only where governments and stakeholders understand the concept of Biosecurity and see the benefits it can be expected to provide will there be the will to make institutional and legislative change.

National consultations will also be essential to build consensus around the most sensitive aspects of any proposed reforms, which will in turn encourage compliance once the reform is undertaken. Moreover, widespread consultations facilitate the circulation of draft legislation among relevant parties, which permits it to be modified in light of inconsistencies with other draft legal instruments being proposed in related areas. All these elements contribute to the design of feasible reforms, suited to the needs and realities of each country wishing to implement a Biosecurity approach.
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