

GOVERNMENT OF THE FEDERAL REPUBLIC OF NIGERIA <u>Application Form for the registration of the gene of Genetically</u> <u>Modified Plant</u>

Ref. No.---

This form should be forwarded to the **Director General/Chief Executive Officer, National Biosafety Management Agency, Plot 393/394 Augustus Aikhomu Way, Utako District, Abuja,** on completion.

1. Administrative information

Purpose of Application:

[(name of Crop species and introduced trait)

Previous Applications or Approvals:

[Information on the status of this crop and trait, including pending, approved, or denied applications for field trials and commercial releases here or in other jurisdictions. Indicate also if this is a new application or a renewal.]

Applicant:

[Name of applying institution/Company, which may also include the name of the key personnel.]

Institutional/Company Address:

Telephone (s):

E-Mail:

Contact Details of Key Person:

Name of Key Person:

Designation of Key Person:

Address:

Telephone (s):

E- mail:

PART I

1. BRIEF DESCRIPTION OF THE GENETICALLY MODIFIED PLANT'S GENE

Include specific and common names of the plant, the country of origin of the plant and a description of the genetically modified trait.

- 2. RELEASE
 - 2.1 Detail specific instructions for the storage and handling of the plant gene, or viable plant parts,
 - 2.2 What type of release is intended,
 - 2.3 When will the release be implemented?
 - 2.4 Where will the release take place?
 - 2.5 detail the type of environment and the geographical areas for which the plant is suited.
 - 2.6 Who will undertake the release?
 - 2.7 Estimate the amount of production of the genetically modified plant within Nigeria per annum, or the amount of viable plant product to be imported into Nigeria per annum.

3. DESCRIPTION OF ANY PRODUCT TO BE DERIVED FROM THE PLANT GENE

- 3.1 identify the part of the plant to be used for the product, the type of product, and the use of the product, the market sector in which the product will be marketed and the trade name of the product.
- 3.2 Specify the exact conditions of use of the product.
- 3.3 provide information on the proposed labeling of the product for marketing
- 3.4 state whether the benefits of the products are available in any other non-genetically modified form. If so, state why the genetically modified form should be approved for I release when other, non-modified products are available.
- 3.5 Detail specific instructions for the storage and handling of plant's gene products that will avoid misuse or escape of the genetically modified plant into an environment for which it was not intended.
- 3.6 detail the likelihood of the genetically modified plant gene or its products being exported from Nigeria, particularly if such export could result in the introduction of the plant's gene into its centre of origin.
- 4. BRIEF SUMMARY OF FIELD TRIALS UNDERTAKEN
 - 4.1 Submit a list of previously authorized activities with the gene in(a) Nigeria
 - (b) other countries
 - 4.2 Include information on the country, year, location and the authority form which permission was obtained to run the field trials.
 - 4.3 Provide full date on the field performance of the genetically modified plant, including the efficacy of the introduced gene trait
- 5. POLLEN SPREAD
 - 5.1 identify all methods of pollination applicable to plant containing the gene

5.2 Identify pollinating agents and the distances to which pollen containing the gene is known to spread.

- 5.3 identify any plants in the are of the release that may become cross-pollinated with the genetically modified pollen.
- 5.4 Describe methods to be used to prevent the spread of genetically modified pollen to wild type plants
- 6. SEED DISPERSAL

6.1 If the seed containing the gene is to be sold, state whether the seed is hybrid

- 6.2 Describe methods to be used to limit the dispersal of genetically modified seed into the environment
- 6.3 If seed dispersal will occur describe what volumes of seed are likely to be dispersed, how this seed will interact in the environment and what long term effects the seed is likely to have on the environment.
- 7 VEGETATIVE SPREAD OF THE GENETICALLY MODIFIED PLANTS
 - 7.1 Describe methods of vegetative reproduction that are available to plant containing the gene
 - 7.2 Describe methods to be used to limit vegetative spread of the plant containing the gene into the environment
- 8. RESISTANCE
 - 8.1 Detail whether the gene is able to initiate resistance, in any biotic components of the environment, to any biologically active foreign gene product.
 - 8.2 Detail what methods are available to minimize the risk of resistance developing in the environment
 - 8.3 Detail how resistance will be managed during release of the plant containing the gene

9. HUMAN AND ANIMAL HEALTH

- 9.1 State whether the plant containing the gene or its products will entre human or animal food chains.
- 9.2 Detail the results of experiments undertaken to determine the toxicity of the gene products to humans and animals
- 9.3 if the gene products are toxic or allergenic in any way, detail how the release will be managed to prevent contact with animals or humans that will lead to discomfort or toxicity.

9.4 What are the implication of the proposed activity with regard to the health and safety of the workers, cleaning personnel and any other person, that will be directly or indirectly involved in the activity?

9.5 Further to the question raised above, indicate the proposed health and safety measures that would be applied to safeguard employees during the proposed activity

10. ENVIRONMENTAL IMPACT AND PROTECTION

- 10.1 Detail any long-term effect the release of the plant containing the gene is likely to have on the biotic and abiotic components of the environment
- 10.2 provide data and information on ecosystems that could be affected by use of the plant containing the gene or its products
- 10.3 Specify what effect the release of the plant containing the gene will have on biodiversity
- 10.4 Specify the measures to be taken in the event of the plant containing the gene or product being misused or escaping into an environment for which it is not intended
- 10.5 If the gene gives rise to crops resistant to agrochemicals, provide information on the registration of the agrochemicals to be used on the crops.
- 11. SOCIO-ECONOMIC IMPACTS Specify what, if any, positive or negative socio-economic impacts that plant containing the gene will have on communities in the proposed region of release if its general release.

- 12. MONITORING AND ACCIDENTS
 - 12.1 indicate the methods and plans for monitoring of the plant containing the gene
 - 12.2 Indicate any emergency procedures that will be applied in the event of an accident

13. PATHOGENIC AND ECOLOGICAL IMPACTS

14.1 Submit an evaluation of the foreseeable impacts, in particular any pathogenic and ecologically disruptive impacts

14. WASTE DISPOSAL

Where only a portion of the gene or plant containing the gene will be used for the product, how will the unused gene or plant parts be disposed of?

15. RISK MANAGEMENT

15.1 Please indicate any risk management measures that would be required during the trial

16. Directions for the applicant:

(This page must should be excluded from the documents submitted to the Director General/CEO's office)

- Please complete all relevant sections f the questionnaire CLEARLY.
- Please provide 5 copies of the application with confidential information.
- Please provide an additional hard copy of the application containing no confidential information. The latter application will be made available for public scrutiny
- Please provide an electronic and hard copy of a risk assessment conducted in accordance with Annex III of the Cartagena Protocol on Biosafety and in the format prescribed below,
- Please submit all relevant documentation to the Director General/Chief Executive Officer at the address indicated in the application form
- The appropriate fee stipulated must paid into the Agency's Bank account after the application has been acknowledged. Please note that the Director General/Chief Executive Officer's office does not accept cash.

17. Declaration

I hereby certify that the information in the application and all attachments is complete and accurate to the best of my knowledge and belief:

Name and Signature of Representative of for Applying Institution/Company:------

Date:

COMMON FORMAT FOR RISK ASSESSMENT

(In accordance with Annex III of the Cartagena Protocol on Biosafety)

	Risk Assessment Details	
1.	Country of taking Decision	Nigeria
2.	Title:	<text entry=""></text>
3.	Contact details	Standard Contact address details: name, function (job title/designation), organization, address, phone, fax, e- main, website
4.	Name and identification of the living modified organism	<text entry=""></text>
5.	Unique identification of the living modified organism:	<text entry=""></text>
6.	Transformation event:	<text entry=""></text>
7.	Introduced or Modified traits:	 Choose the trait from the following list: A. Abiotic Environmental tolerance ✓ Altered photoperiod sensitivity ✓ Cold or heat tolerance ✓ Drought or water tolerance ✓ Other abiotic environmental tolerance B. Altered Growth, Development and product quality ✓ Altered ripening or flowering ✓ Coloration ✓ Fertility ✓ Growth rate or yield ✓ Male sterility ✓ Other growth, development and product quality ✓ Other growth, development and cinc.allergenicity ✓ Other growth, development and product quality ✓ Selected marker genes and reporter gens ✓ Uptake or degradation of environment pollutants

		 Chemical Tolerance Herbicide tolerance Other chemical tolerance Medical Products Animal vaccines Development of transplant organs Other medical products Production of pharmaceuticals Pest Resistance Bacterial resistance Fungus resistance Insect resistance Nematode resistance Other pest resistance Virus resistance and <text entry="" for="" list="" not="" on="" other,="" the=""></text>
8.	Techniques used for modification:	<controlled common<br="" for="" vocabulary="">techniques – please select techniques used for the transformation: plasmid carried by <i>agrobacterium tumefacients</i>, biolistic methods, breeding, electric shock (poration), asmotic shock> and <text entry="" for="" list="" not="" on="" other,="" the="" –=""></text></controlled>
9.	Description of gene modification:	<text entry=""></text>
		Characteristics of modification
10.	Vector characteristics (Annex III.9(c)):	<text and="" any,="" characteristics="" entry="" host="" identity,="" if="" include="" its="" of="" or="" origin,="" range<="" should="" sources="" th="" the="" vector,="" –=""></text>
11.	Insert or inserts (Annex III.9(d)):	<text -="" characteristics="" entry="" genetic="" of<br="">the inserted nucleic acid and the function it specifies, and/or characteristics of the modification introduced></text>
		Recipient organisms or parental organisms (Annex III.9(a)):
12.	Taxonomic name/status or recipient organisms or parental organisms:	<pre><controlled agreed="" international="" standards="" vocabulary:=""> and <text entry=""> - for other, not on the list</text></controlled></pre>
13.	Common name of recipient organism or parental organisms	<controlled thesaurus="" vocabulary="" with=""> and <text entry="" for="" not="" on="" other="" the<br="" –="">list></text></controlled>
14.	Point of collection or acquisition of recipient or parental organisms:	<text entry=""></text>

15.	Characteristics of recipient	<text entry=""></text>
	organism or parental organisms:	
16.	Centre(s) of origin of recipient organism or parental organisms	<text describe="" entry="" exact="" location<br="" the="" –="">and give geographical coordinates></text>
17.	Centres of genetic diversity, if known of recipient organism or parental organism may persist or proliferate	<text describe="" entry="" exact="" location<br="" the="" –="">and geographical coordinates></text>
18.	Habitats where the recipient organism or parental organisms may persist of proliferate: Donor organism or organism (Annex III.9(B)):	<text description="" entry="" habitat<br="" of="" the="" –="">where the organism may persist or proliferate></text>
19.	Taxonomic name/status of donor organism(s)	<controlled agreed<br="" vocabulary:="">international standards> and <text entry<br="">for other, not on the list></text></controlled>
20.	Common name of donor organism(s):	<controlled thesaurus="" vocabulary="" with=""> and <text entry="" for="" list="" not="" on="" other,="" the=""></text></controlled>
21.	Point of collection or acquisition of donor organism(s):	<text and="" coordinates="" entry="" exact="" geographical="" location="" the="" –=""></text>
22.	Characteristics of donor organism(s) related to biosafety: Intended use and receiving environment	<text biological<br="" entry="" relevant="" –="">characteristics of donor organisms></text>
23.	Intended use of the LMO (Annex III 9(g)):	<text entry="" information="" relating="" the<br="" to="" –="">intended use of the living modified organism, including new or changed use compared to the recipient organism or parental organisms></text>
24.	Receiving environment (annex III.9(h)):	<text entry="" information="" on="" the<br="" –="">location, geographical, climatic and ecological characteristics, including relevant information on biological diversity and centres of origin of the likely potential receiving environment></text>
	Risk Assessment Summary	
25.	Detection/identification method of the LMO (annex III.9(f)):	<text and<br="" detection="" entry="" suggested="" –="">identification methods and their specificity, sensitivity and reliability></text>
26.	Evaluation of the consequences (Annex III.8(b)	<text an="" entry="" evaluation="" of="" the<br="" –="">likelihood of these adverse effects being realized, taking into account the level and kind of exposure of the likely potential receiving environment to the living modified organism></text>

27.	Evaluation of the consequences	Text entry – An evaluation of the
	(Annex III.8(c)):	consequences should these adverse
		effects be realized>
28.	Overall risk (annex III.8(d))	<text an="" entry="" estimation="" of="" overall<="" th="" the="" –=""></text>
		risk posed by the living modified
		organism based on the evaluation of the
		likelihood and consequences of the
		identified adverse effects being realized>
29.	Recommendation (Annex III.8(e)):	<text a="" as="" entry="" recommendation="" th="" to<="" –=""></text>
		whether or not the risks are acceptable or
		manageable, including, where necessary
		identification of strategies to manage
		these risks>
30.	Actions to address uncertainty	<text about="" any="" details="" entry="" further<="" th="" –=""></text>
	regarding the level of risk (annex	information that has been requested
	III.8(f)):	where there is uncertainty regarding the
		level of risk as well as any information on
		risk management strategies and/or
		monitoring of the LMO in the receiving
		environment>
31.	Availability of detailed risk	<text entry="" indicate="" please="" th="" whether<="" –=""></text>
	assessment information:	more details on the irks assessment are
		available and how they can be accessed >
32.	Any other relevant information:	<text any="" entry="" information="" other="" th="" that<="" –=""></text>
		is relevant to the risk assessment. E.g.
		information of non-CBI nature that was
		included in the original application but is
		not included in this form>
33.	Attach document:	Not applicable to applicant
		<specific entry:="" of="" option="" th="" to<="" types=""></specific>
		choose a file form the local source and
		'upload' a copy of the BCH server>
34.	Notes:	<text only=""></text>