Form20-1	IASRI Safety	Office←Person	in charge of	managing	experiments←Project Leader
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Safety Supervisor	Confirmation by Safety Office	Receipt by the Safety Office	20YY, MM, DD
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Application Form for Genetic Modification Experiment

Date of submission: June 1, 2016

To: The Director General, Japan Synchrotron Radiation Research Institute

(Person in charge of experiment) ¹⁾	
Name of the organization	
XX University, Graduate School	
Department and title	
XX Department XX Assistant	
Name (print and signature)	
Koukido Hanako	
(Manager) ²⁾	
Name (Leave it blank if User is applying) Seal	

To carry out the following genetic modification experiment, I hereby apply for approval of the genetic recombination committee.

Receipt number ³⁾	(Do not write when submitting) Prolongation of duration due to the completion of the valid period of the approved experiment → "Renewal"					
	□ New Change in experimental material or place → "Amendment" Check appropriate box and write the previous receipt number.					
Type of application ⁴⁾	☐ Renewal (Previous receipt num Check appropriate box and write the previous receipt number.					
	Amendment (Previous receipt number Approval 03-10)					
Title of experiment ⁵⁾	Examination of metabolism of XXX enzyme in XX deficient mice.					
Title of experiment	The title should concisely describe the purpose and a summary of the experiment					
	■ Microbiology experiment					
Type of experiment ⁶⁾	☐ Large-scale cultivation experiment					
Type of experiment	■ Animal experiment (■ Animal inoculation · □ Animal modification)					
	☐ Plant experiment (☐ Plant inoculation • ☐ Plant modification • ☐ Fungus modification)					
	The dynamics of XXX enzyme will be examined by infecting mice with a virus that expresses					
The purpose	XXX enzyme due to a mutated XX gene.					
	The flow of the relevant genetic modification experiment should be stated.					
Summary ⁷⁾						
,	The validity period of the experiment is a maximum of three years from the					
	The validity period of the experiment is a maximum of three years from the date of approval. Enter the desirable start date and planned completion date.					
Expected dyration of	October **, 2016 to March 31, 2019					
Expected duration of experiment ⁸⁾	October 11, 2010 to March 31, 2019					
схренниен	Address (Postal code)					
Contact information of	1-1-1 Koto, Mikazuki-cho, Sayo-gun, Hyogo					
the person in charge of	Phone (ext./PHS) 07**—***(2***)					
the experiment	Fax: 07**-**-					
r r r	E-mail: hanako@*******.ip					

Form20-1 JASRI Safety Office←Person in charge of managing experiments←Project Leader

	Organization and department of the person in charge of communications
	XX University, Graduate School XX Faculty XX Lab Office
	Name of the person in charge of communications
	Last name First name
0))	Address (Postal code)
Other contacts ⁹⁾⁾	1-1-1 Koto, Mikazuki-cho, Sayo-gun, Hyogo
	Phone (ext./PHS) 07**—**—****(3***)
	Fax: 07**-**
	E-mail: shikaku@*******.**.jp

Place of the experiment (place	ce for animal keeping and raising) ¹⁰⁾ ar	nd place	for storag	e of gene	tically n	nodified o	organism	ns ¹¹⁾⁾
Building	Room	Containment measures						
Dunding		P1	P1A	P1P	P2	P2A	P2P	Storage
	□BL20B2 experimental hutch				<u></u>			
	□BL28B2 optics hutch							
	□BL40XU experimental hutch							
77 1 177 11	□BL20B2 animal operation room	Y 1 1						
Experiment Hall	☐Mobile operation room	I I I						
	□BL41XU experimental hutch	1 I I						
	□BL38B1 experimental hutch	•						
	□BL32XU experimental hutch	7 1 1						
	■ Mouse room	1 1 1 1	0					
Experimental Animal	Genetic experiment room	I I I	0					
Facility	☐Treatment room							
	□BL20B2 experimental hutch							
Medium-length Beamline Facility (Experiment building)	□BL20XU experimental hutch	•						
	☐Animal operation room	T						
(Experiment building)		Y				-	• • • • • • • • • • • • • • • • • • •	
	□Room 101	î !						
	□Room 201	1 1 1						
	□Room 202	I I I						
Medium-length Beamline	□Room 204	# I I					<u>.</u>	
Facility	□Room 212	7 1 1						
(Research building)	□Room 213	Y						
	■Biochemistry lab 1 Room 208	Attach	documents	explaining t	the follow	ings.		
	☐Biochemistry lab 2 Room 209	(1) Plac drawing	documents ce of major g of the exp	facility, equ erimental a	ipment, a ırea)	ind instrum	ent (a det	ailed
	☐Biochemistry lab 3 Room 210	(2) The	state of an	ımals or pla	ints that a	are kept in t	he room (of area,
	□Experimental hutch (EH3)	Date dire	7			1		
SACLA	☐Biological sample prep. room							
Others	***** building room	I I I	0					
(Write the name of the building and room)		 						
bulluing and foom)	 	I I I						

Nucleic acid donor	Experiment	Donor nucleic acid (type of nucleic	Identification	Note
	classification	acid)		
(Target gene)			Completed/Not completed	
Streptomyces lavendulae ##	Class 1	XX enzyme gene (cDNA)	Completed	
strain: genus Streptomyces				
bacteria of the family				
Streptomycetaceae				

Form20-1 JASRI Safety Office←Person in charge of managing experiments←Project Leader

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Protostome:Schistosoma	Class 2	Glutathione S-transferase (cDNA)	Completed		
japonicum					
When changing experim	ental materials, und	erline the materials to be added.			
		J	Completed		
Mouse	Class 1	XX enzyme gene (cDNA)	Completed		
(Expression regulatory gene)					
E. coli of the family	Class 1	tac promoter (genome DNA)	Completed		
Enterobacteriaceae		(Section 2 : 1.2)			
E. coli of the family	Class 1	lac promoter (genome DNA)	Completed		
Enterobacteriaceae		, ,			
Bacteriophage T7	Class 2	T7 promoter (genome DNA)	Completed		
E. coli of the family	Class 1	Ampicillin-resistant gene	Completed		
Enterobacteriaceae		(genome DNA)			
(Selectable marker gene)					
Host/vector ¹³⁾					
Host	Experiment	Vector	Type	Note	
	classification	v Cotton	1,700	11010	
			Microorganism/Animal/Plant		
EK1	Class 1	pUC119 (cloning vector for E. coli)	Microorganism		
		pGEX (vector for protein expression)			
SD rats and fertilized eggs of	Class 1	Not used	Animal		
SD rats					
Characteristics of animals,		be used for cultured cells that will be			
plants, or cells that possess		The cultured cell line that will be use			
genetically modified	•	than on the artificial medium. The gen		•	
organisms. 14)		will be expressed by infection. The in		to virus	
Table County II I'm	multiplication.	The infected cells will not develop dr	ug resistance.		
Table of genetically modified	og nom offereler 1				
organisms and containment measures. 15)	as per attached				
Method of inactivation of	Inactivate by o	utoclaving (121°C, 20 minutes).			
genetically modified	machivate by at	atociaving (121 C, 20 initiates).			
organisms. 16)					
Note ¹⁷⁾	Grant-in-Aid fo	or Scientific Research of Japan Synchi	rotron Radiation Resear	ch Institute	
	will not be used.				
		g (such as Grant in Aid for Scientific	Research of the Ministr	y of Education,	
		, Science and Technology) is received	l as Japan Synchrotron F	Radiation	
	Research Institu	ute, indicate it accordingly.			

XNotes

- 1) For "Project Leader," the information shall be given on a person who is directly manage a genetic recombination experiment at SPring-8 and have experience for one year or more. However, a student should not be.
- 2) For "Person in charge of managing experiments," the information shall be given on a person who is in charge of administration of this application.
 - JASRI, RIKEN, JAERI Staff >> Director
 - User >> Director of Users Office (A blank is sufficient in case it submits.)
- 3) Since "Proposal number" is informed when Safety Office receives this application, leave a column blank. Proposal number is required for all the documents for which you will apply in the future.
- 4) For "Type of Application," select any items under which your application falls. In case Continuation or Changes, give the previous proposal number.
- 5) For "Title," mention a name that expresses the objective and an outline of a genetic recombinant experiment briefly.
- 6) For "Type," select all items under which a genetic recombinant experiment falls.
- 7) For "Outline," all living modified organisms involved in a genetic recombinant experiment and the categories of containment measures to be taken during a genetic recombinant experiment shall be mentioned so as to show their processes.
- 8) The experiment is valid for a maximum of three years from approved day.
- 9) For "Other contact," if there is any other contact for administrative matters than Project Leader or Deputy Project Leader, give the information on the contact.
- 10) For "Laboratory Experiment area Experiment section (include the area of the breeding animals or culture of plants)," select all area under which a genetic recombinant experiment falls. If there is no appropriate column, indicate name of facility and room on proper column and attach the information those mentioned in below.
 - ①Names and positions of major facilities, equipment and apparatus;
 - ②In case an animal or a plant that has no relation with a genetic recombination experiment is bred or cultured in the laboratory, the experiment section, the experiment area, the breeding section or the screened greenhouse, the state of the breeding of the animal or the culture of the plant;
- 11) For "Facility of Storage," select all area that storage living modified organisms in the process of a genetic recombination experiment. If there is no appropriate column, indicate name of facility and room on proper column and attach the information about them.
- 12) For "Donor organism/Donor nucleic acid," the following shall be mentioned about the donor organism and donor nucleic acid of the living modified organism for Genetic recombination experiment every component elements (target genes, expression regulatory genes, drug-resistant genes and marker genes).
 - a. General name and taxonomical position (familia, genus, species, strain) of donor organism
 - b. General name and type (such as genomic nucleic acid, complementary deoxyribonucleic acid or synthesized nucleic acid) of donor nucleic acid.
 - c. Attach copy of nucleotide sequence information or an accession number to the nucleotide sequence database of, for example, the Japan DNA Databank (only in the case of donor nucleic acid that is identified nucleic acid).
- 13) For "Recipient organism/Vector," the following shall be mentioned about the recipient organism and vector of the living modified organism for Genetic recombination experiment.
 - a. General name and taxonomical position (familia, genus, species, strain) of recipient organism
 - b. General name, code and short explanation about (ex. pUC119 cloning vector for *E.coli*)
 - c For "Type," select Microorganisms, Animals or Plants.
- 14) For "Characteristics of animal, plant or cell which retains living modified organisms," in addition to items those mentioned in below, characters expected to be newly given or already given to an animal, a plant or a cell which retains the living modified organism for Genetic recombination experiment in comparison with animals, plants or cells which do not retain the living modified organisms Genetic recombination experiment shall be mentioned.
 - a. Taxonomical position and experiment classification of animal, plant or cell which retains living modified organism;
 - b. State of distribution in natural environment and environment in which living or growth is possible;
 - c. Pathogenicity, production of harmful substances and other properties;
- 15) For "Combination Living Modified Organism and its Category of Containment Measures," all donor organisms, donor nucleic acids, vectors, recipient organisms and animals, plants or cells which retains living modified organism involved in a genetic recombination experiment and the categories of containment measures to be taken during the experiment shall be mentioned so as to show processes of the experiment.
- 16) For "Measure for inactivating living modified organism," about the containment measures to be taken during Genetic recombinant experiment, mention a measure for inactivating waste products containing the living modified organism and apparatus and appliances to which the living modified organism sticks, and the effectiveness of the measure.
- 17) When you have received grant of public costs from JASRI, give a note.

Write the combination of nucleic acid donor, donor nucleic acid, vector, host, and possessing organisms along with the containment measures taken in the relevant step of the experiment so that the flow of the experiment is clear. When changing experimental materials, underline the materials to be added.

Nucleic acid donor	Donor nucleic acid	Vector	Host etc.	Possessing organisms	Containment Measure Classification	Note
Streptomyces lavendulae ## strain Mouse E. coli	XX enzyme gene (cDNA) XX enzyme gene (cDNA) lac promoter (genome DNA) Ampicillin-resistant gene	pUC119	JM109 derived from E. coli K12 strain	None	P1	Level B1 Cloning
Mouse Bacteriophage T7 E. coli	XX enzyme gene (cDNA) T7 promoter (genome DNA) Ampicillin-resistant gene lac I (genome DNA)	pET	BL21 (DE3) derived from E. coli B strain	None	P1 Indicate the step of operation.	Level B1 Protein expression
E. coli Schistosoma japonicum Streptomyces lavendulae ## strain	tac promoter (genome DNA) lac I (genome DNA) Glutathione S-transferase(cDNA) XX enzyme gene (cDNA)	pGEX	BL21 derived from E. coli B strain JM109 derived from E. coli K12 strain	None	P1	Level B1 Protein expression