



Risk Assessment for Genetically Modified Micro-organisms [GMMs]

### Class 1 & Class 2

The following constitutes an **example** of a Risk Assessment for a GMM falling within Class 2. It is designed as a guide to how a Risk Assessment should be conducted, the elements which should be considered during a Risk Assessment (the recipient organism, the insert, the vector, the host, the final GMM, human health and environmental considerations, containment measures) identification of possible harmful effects and the possibility of those effects occurring. The Risk Assessment process culminates in the determination of the final classification of the GMM and the appropriateness of the corresponding containment measures.

### **Description of each GMO:**

Recombinant Adenovirus viral vectors containing a gene encoding a therapeutic protein VEGF - which promotes the growth of vascular endothelial tissue.

#### Purpose of the contained use:

The aim is to produce live intact Adenovirus viral vectors. The adenovirus vector contains a gene which codes for human vascular endothelial growth factor. This protein has the potential to be effective in the treatment of coronary artery disease and peripheral vascular disease.

## PROCEDURE 1

(3.1) Identification of harmful properties (hazard) of the GMM:	
(3.2.1) the recipient	The recipient organism is human Adenovirus type 5. It is
organism	ubiquitous, causes only a mild respiratory disease in humans
	which is self limiting and does not require any specific
	treatment. Similarly there is no association with allergic or
	toxic effects. The complete E1 region and the majority of the
	E3 region of the genome have been removed thereby





	rendering the viral vector replication deficient.
(3.2.2) the insert	The genetic insert (VEGF) is human in origin and poses no risk
	to animal health or the environment.
(3.2.3) the vector	Construction of recombinant Adenovirus is a two step
	process in which the desired expression cassette is first
	assembled into a pUC vector and subsequently transferred
	into the Adenoviral genome by homologous recombination.
	pUC vectors have a history of safe use. The Adenovirus viral
	vectors are replication defective by virtue of deletion of the
	E1 and E3 regions.
(3.2.4) the host	The host cell is PER.C6 cell line which is derived from human
	embryonic retinoblasts transformed with the E1 region of
	Adenovirus 5. Since the Adenoviral vector is replication
	deficient recombinant Adenovirus can only grow in
	complementing cells such as PER.C6 which contain the
	appropriate E1 sequences.
	PER.C6 cells die rapidly outside the artificial environment
	created within the laboratory. There is little likelihood of the
	recombinant PER.C6 cells proliferating or surviving in the
	environment and therefore poses little risk to animal/plant
	health or the environment
(3.2.5) the resulting GMM	Recombinant Adenovirus vector contains the gene for human
	vascular endothelial growth factor. The recombinant
	Adenovirus is replication deficient and therefore can only
	replicate in cells which carry complementing regions of the
	E1 genes. It will not replicate in other in vivo or in vitro cells.
	Furthermore since the modified virus is replication deficient
	it is less pathogenic than the wild type and there is minimal
	capacity for colonisation. If it is exposed to the environment
	it is unlikely to survive for extended periods.





	A replication competent adenovirus has the potential to be
	produced however this is unlikely given that the E1
	sequences in the PER.C6 cells do not overlap with the
	deleted E1 region. Therefore in order to generate a
	replication competent adenovirus two non homologous
	recombination events would have to occur. A revertant
	regaining the E1 gene would still be devoid of the E3 gene
	since PER.C6 does not contain the E3 gene. The absence of
	the E3 gene would reduce the fitness of the virus as an
	infective agent.
(3.2.5.1) Human	Wild type human Adenovirus type 5 is ubiquitous and causes
Health Considerations	self-limiting infections of the upper respiratory tract and the
considerations	common cold. There may be a possibility that the modified
	adenovirus may mimic some of the characteristics of the
	wild-type however recombinant Adenovirus can only
	replicate in complementing cells such as PER.C6. Even if
	replication competent Adenovirus 5 were generated the risk
	associated is low since human adenoviral infection is very
	common and the majority of adults have already been
	infected .
(3.2.5.2)	There may be a possibility that the modified adenovirus may
Environmental Considerations	mimic some of the characteristics of the wild-type. However
considerations	recombinant Adenovirus can only replicate in
	complementing cells such as PER.C6 and consequently poses
	a low risk to animal/plant health and the environment.
	Level 2 containment measures will be in operation and the
	principles of Good Microbiological Practice will be applied. In
	conjunction with this access to the laboratory will be
	restricted when work with infectious agents is in progress.





infection may have serious consequences will not be allowed to enter the laboratory. A biohazard sign will be posted at the laboratory entrance bearing appropriate information including the agent(s) in use, containment level, the investigator's name and telephone number, personal protective equipment requirements and exiting procedures if any. Biosafety procedures will be incorporated into Standard Operating Procedures or the biosafety manual and personnel will be advised of special hazards. All work will be done with the approval of the safety sub-committee.

### (3.3) Initial classification of the GMM:

Human Adenovirus Type 5 is ubiquitous, causes only a mild respiratory disease in humans which is self limiting and does not require any specific treatment. The Adenovirus viral vectors are replication defective owing to the removal of the E1 and the E3 regions. The genetic insert (VEGF) is human in origin and poses no risk to animal/plant health or the environment. The pUC vectors have a history of safe use. Modified adenovirus may mimic some of the characteristics of the wild-type or may pose a risk to immuno compromised individuals. It is therefore allocated to Class 2, GMM activities of low risk for which level 2 containment is appropriate to protect human health as well as the environment.

(3.4) Assessment of possibility of harmful effects occurring:		
(3.4.1) Nature of	The experiments entail standard molecular biology	
activities to be	techniques which carry no special risk and all are	
undertaken	conducted in accordance with safety SOPs held in house.	
(3.4.2)	A typical viral preparation involves propagation in 30	
Concentration and	flasks each containing 30ml of culture medium The cell	
scale	pellet is harvested from these flasks and the virus is	
	released by repeated freeze thawing. The virus is isolated	
	and a typical yield is 1 x 1010 pfu/ml	
(3.4.3) Culture	Adenoviral vectors will be propagated in PER.C6 cells. Culture	
conditions		
	conditions are as per 3.4.2 above. The GMM is incubated at	
	32°C for 36 – 48 hours.	





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(3.4.3.1)	Only the immediate laboratory environment is likely to be
Environment likely to be exposed	exposed and as outlined in section 3.2.5.2 considerable care
	is taken to ensure that the GMM is contained and that
	areas/personnel beyond the laboratory are not exposed to
	the GMM. Furthermore it is expected that the GMM will not
	survive for significant periods in the environment.
(3.4.3.2) Presence	Neither animals nor plants are susceptible to human
of susceptible species	Adenovirus Type 5. Laboratory staff and/or immuno
	compromised persons constitute those most at risk and as
	outlined in section 3.2.5.2 considerable care is taken to
	ensure that the GMM is contained.
(3.4.3.3) Whether	Recombinant Adenovirus is replication incompetent by virtue
the environment can support the	of the fact that the E1 and the E3 gene sequences have been
survival of the	removed. It is therefore only capable of replication in
GMM	complementing cells such as PER.C6.
(3.4.3.4) Effects on	Since the GMM is considered incapable of survival in the
the physical environment	environment no effects on the physical environment are
	expected.

# PROCEDURE 2





(4.1) Determination of final classification and containment measures Wild type human Adenovirus type 5 is classified as Class 2. It is ubiquitous, causes only a mild respiratory disease in humans which is self limiting and does not require any specific treatment. There is no association with allergic/toxic effects. The genetic insert (VEGF) is human in origin and poses no risk to animal/plant health or the environment. pUC vectors have a history of safe use and Adenoviral vectors are replication defective by virtue of deletion of the E1 region. Modified adenovirus may mimic some of the characteristics of the wild-type virus or may pose a risk to immuno compromised persons. It is therefore allocated to Class 2, GMM activities of low risk for which level 2 containment is appropriate to protect human health as well as the environment

The GMM activity is therefore classified as Class 2.

### (4.2) Confirmation of adequacy of final containment measures

Principles of Good Microbiological Practice and Good Occupational Safety and Hygiene in accordance with Part A of the Fourth Schedule of S.I. No. 73 of 2001. The requirements of Containment Level 2 as given in table 1A - 'containment measures for contained use of GMOs in a laboratory' - of S.I. No. 73 of 2001.