



# Biological Risk Group and Containment Level Assessment

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This template is provided by the Public Health Agency of Canada (PHAC) and serves as a tool to determine the Risk Group (RG) and Containment Level (CL) requirements of organisms and biological materials for teaching, research, or other activities at the University of Lethbridge.

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## INSTRUCTIONS

1. If the organism, biological material or biohazardous material risk group and containment level is not known, a complete biological risk group and containment level assessment must be performed. Complete all applicable sections of PHAC’s risk assessment template on the following pages.
  2. The Biosafety Officer is available to assist individuals with the completion of the Biological Risk Group and Containment Level Assessment.
  3. Submit this form to the [Biosafety Officer](#) when completed. This document is to be submitted with the Biosafety Plan.
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# Biological Risk Group and Containment Level Assessment

Performed by:  
Date:

## Pathogen Risk Assessment

Pathogen Name:

Taxonomy:

- Kingdom:
- Family:
- Subfamily:
- Genus:
- Species:
- Sub-Species:
- Other (e.g., clonal isolate, serotype, serovar, biovar):

### 1. Pathogen Oversight

#### Regulatory Authorities

During your literature search, determine whether the agent has the ability to infect humans, terrestrial animals, aquatic animals, plants or bees. Even opportunistic infections should be noted, regardless of the risk group outcome of your assessment. Identify whether the agent is subject to official control. This will help you identify who you will need to contact in order to work with the agent in the laboratory.

Is the pathogen a

- ☐ strain, clonal isolate, or recombinant variant of a pathogen with a known risk group (RG)?
- ☐ human pathogen?\*
- ☐ terrestrial animal pathogen?\*
- ☐ non-indigenous animal pathogen?\*\*
- Notes:
  - ☐ [OIE listed disease](#)?\*\*
- Notes:
- ☐ aquatic animal pathogen?\*\*
- ☐ plant pathogen?\*\*
- ☐ bee pathogen?\*\*

**\*Human and terrestrial animal pathogens may be regulated by the Public Health Agency of Canada.**  
**\*\*Terrestrial animal pathogens that are non-indigenous to Canada (cause foreign animal and emerging animal diseases), aquatic animal pathogens, plant pathogens, and bee pathogens may be regulated by the Canadian Food Inspection Agency.**

Is the pathogen subject to official control?

- ☐ [National Notifiable Disease](#)
- ☐ [Domestic Substances List](#)
- ☐ [Reportable Disease](#)
- ☐ [Immediately Notifiable Disease](#)
- ☐ [Annually Notifiable Disease](#)
- ☐ [Plant Protection Regulations](#)
- ☐ [Quarantine Act](#)
- ☐ Provincial Notifiable Disease
- Other (list):

#### Biosecurity Oversight

**Biosecurity** refers to security measures designed to prevent the loss, theft, misuse, diversion, or intentional release of pathogens, toxins, and other related assets (e.g., personnel, equipment, non-infectious material, and animals).

- Identify whether the agent appears on any of the lists of agents of potential concern for biosecurity. Agents on these lists may be subject to additional security requirements.
- Identify whether there are any biosecurity considerations that should be noted in the risk assessment. Provide a brief summary, supported by references where possible, as to the potential biosecurity concerns related to this agent. Any biosecurity concerns should be fully elaborated in your [Biosecurity Risk Assessment](#) and [Biosecurity Plan](#).

- ☐ [Australia Group Common Controls List](#)
- ☐ [Select Agents and Toxins List](#)
- ☐ [Security Sensitive Biological Agent](#)
- ☐ This pathogen has no known biosecurity concerns

Notes (full details should be elaborated in your Biosecurity Plan):

### 2. Pathogen Description

Provide background information that could be relevant to the interpretation of the risk assessment or overall risk. Provide references to support your comments. Some of the types of information that may be applicable to the pathogen risk assessment are listed below.

- **Example 1**, when assessing a recombinant virus, the genome structure of the native virus and modifications should be described in sufficient detail to determine how the modifications will impact the different factors being assessed (e.g., pathogenicity).
- **Example 2**, when assessing bacteria or fungi, the ability to product toxins may directly impact pathogenicity.
- **Example 3**, when assessing fungi with complex taxonomy or numerous changes to taxonomy, current and historical nomenclature should be described.

**Reconstructed, Engineered or modified pathogens** should be assessed throughout the risk assessment by comparing the newly created pathogen to the wild type or a previously assessed variant, linking the various modifications to anticipated effects on the different risk factors (e.g., pathogenicity, communicability).

#### General Information

- Taxonomy
- Historical background
- Size
- Shape
- Structure
- Genome structure/information
- Ideal growth conditions
- Modifications (e.g., CRISPR gene drives)
- Temperature tolerance

#### Bacteria

- Motility
- Sporulation
- Toxin production
- Oxygen requirements
- Gram staining, AF staining
- Enzymatic activity

#### Viruses

- RNA/DNA virus
- Single/Double stranded
- Other classifications

#### Other (e.g., Fungi, Protozoa)

- Life cycle
- Reproduction
- Morphology
- Growth and physiology
- Toxin production



# Biological Risk Group and Containment Level Assessment

## 3. Pathogenicity (Individual Risk)

### Assessment of Human Pathogenicity Indicators

Assess the indicator questions and use these to rate the likelihood of serious disease. Use the rationale section under each question to substantiate your analysis with a description and corresponding references.

Outline uncertainty and assumptions within the rationale for each indicator. The greater the assumptions/uncertainty, the more frequently the risk assessment should be reviewed.

<p>1) If exposed, what is the likelihood that infection would result, with or without overt signs of disease?</p> <p><input type="checkbox"/> None      <input type="checkbox"/> Low      <input type="checkbox"/> Moderate      <input type="checkbox"/> High      <input type="checkbox"/> Unknown</p> <p>Rationale:</p>
<p>2) If exposure led to disease, what is the likelihood that acute signs of disease would be evident?</p> <p><input type="checkbox"/> None      <input type="checkbox"/> Exclusively in susceptible populations      <input type="checkbox"/> Low      <input type="checkbox"/> Moderate      <input type="checkbox"/> High      <input type="checkbox"/> Unknown</p> <p>Rationale:</p>
<p>3) If exposure led to disease, what is the likelihood that there would be serious sequelae or mortality?</p> <p><input type="checkbox"/> None      <input type="checkbox"/> Exclusively in susceptible populations      <input type="checkbox"/> Low      <input type="checkbox"/> Moderate      <input type="checkbox"/> High      <input type="checkbox"/> Unknown</p> <p>Rationale:</p>
<p>4) Are certain populations (e.g., pregnant, elderly, immunocompromised) at an increased risk of infection or disease?</p> <p><input type="checkbox"/> Yes      <input type="checkbox"/> No      <input type="checkbox"/> Unknown</p> <p>Rationale:</p>
<p>Rate the likelihood of serious disease considering the <b>Human Pathogenicity Indicators</b> above.</p> <p><input type="checkbox"/> None, the agent is not a human pathogen;</p> <p><input type="checkbox"/> Low, the agent is an extremely rare opportunistic pathogen. Serious disease may occur in severely ill or immunocompromised;</p> <p><input type="checkbox"/> Moderate, the agent is able to cause serious disease but is unlikely to do; or</p> <p><input type="checkbox"/> High, the agent is likely to cause serious disease.</p>



# Biological Risk Group and Containment Level Assessment

## Assessment of Natural Animal Host(s) Pathogenicity Indicators

Assess the indicator questions and use these to rate the likelihood of serious disease in the natural animal host. **Natural animal hosts** are those where infection and/or disease in the animal would occur in a natural environment, and includes wild animal species (e.g., wild rodents, ruminants, etc.). Information obtained under experimental conditions designed to reproduce natural exposure may also be of use. Other information obtained from experimentally infected animals should be considered as surrogate data only. Use the rationale section under each question to substantiate your analysis with a description and corresponding references.

Outline uncertainty and assumptions within the rationale for each indicator. The greater the assumptions/uncertainty, the more frequently the risk assessment should be reviewed.

1) If exposed, what is the likelihood that infection would result, with or without overt signs of disease? <input type="checkbox"/> None <input type="checkbox"/> Low <input type="checkbox"/> Moderate <input type="checkbox"/> High <input type="checkbox"/> Unknown  Rationale:
2) If exposure led to disease, what is the likelihood that acute signs of disease would be evident? <input type="checkbox"/> None <input type="checkbox"/> Exclusively in susceptible populations <input type="checkbox"/> Low <input type="checkbox"/> Moderate <input type="checkbox"/> High <input type="checkbox"/> Unknown  Rationale:
3) If exposure led to disease, what is the likelihood that there would be serious sequelae or mortality? <input type="checkbox"/> None <input type="checkbox"/> Exclusively in susceptible populations <input type="checkbox"/> Low <input type="checkbox"/> Moderate <input type="checkbox"/> High <input type="checkbox"/> Unknown  Rationale:
4) Are certain populations at an increased risk of infection or disease? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown  Rationale:
Rate the likelihood of serious disease considering the <b>Natural Animal Host Pathogenicity Indicators</b> above. <input type="checkbox"/> None, the agent is not a animal pathogen; <input type="checkbox"/> Low, the agent is an extremely rare opportunistic pathogen. Serious disease ;may occur in severely ill or immunocompromised; <input type="checkbox"/> Moderate, the agent is able to cause serious disease but is unlikely to do so; or <input type="checkbox"/> High, the agent is likely to cause serious disease.



# Biological Risk Group and Containment Level Assessment

## 4. Pre- and Post-Exposure Measures (Human Community Risk)

### Assessment Human Pre- and Post-Exposure Measures Indicators

Assess the indicator questions and use these to rate the level of protection from infection and/or the development of disease. Use the Rationale section under each question to substantiate your analysis with a description and corresponding references.

<div>1) Are pre-exposure measures available to prevent infection or disease (e.g., vaccines, pre-exposure prophylaxis)?<div><input type="checkbox"/> Not available</div><div><input type="checkbox"/> Limited availability</div><div><input type="checkbox"/> Readily available for use on-demand</div><div><input type="checkbox"/> Widely available and in use in the community</div><div><input type="checkbox"/> Unknown</div></div> <div>Rationale:</div>
<div>2) Are these pre-exposure measures effective at preventing infection or disease?<div><input type="checkbox"/> Not applicable, pre-exposure measures are not available</div><div><input type="checkbox"/> Not effective, minimal protection</div><div><input type="checkbox"/> Moderately effective, partial protection</div><div><input type="checkbox"/> Highly effective*, almost complete protection</div><div><input type="checkbox"/> Unknown</div></div> <div>Rationale:</div>
<div>3) Are post-exposure measures available to treat infection or prevent disease (e.g., post-exposure prophylaxis, antibiotics, antifungals, antivirals)?<div><input type="checkbox"/> Not available</div><div><input type="checkbox"/> Limited availability</div><div><input type="checkbox"/> Readily available for use on-demand</div><div><input type="checkbox"/> Widely available and in use in the community</div><div><input type="checkbox"/> Unknown</div></div> <div>Rationale:</div>
<div>4) Are these post-exposure measures effective at treating infection or preventing disease?<div><input type="checkbox"/> Not applicable, post-exposure measures are not available</div><div><input type="checkbox"/> Not effective</div><div><input type="checkbox"/> Moderately effective</div><div><input type="checkbox"/> Very effective</div><div><input type="checkbox"/> Unknown</div></div> <div>Rationale:</div>
<div>5) Are there sub-populations in which the use of or access to pre-exposure measures is less than the general population?<div><input type="checkbox"/> Yes    <input type="checkbox"/> No    <input type="checkbox"/> Unknown</div></div> <div>Rationale:</div>
<div>Rate the level of protection from infection and/or the development of disease considering the <b>Pre- and Post-Exposure Measures Indicators</b> above.<div><input type="checkbox"/> None, if exposed, the community would not be protected;</div><div><input type="checkbox"/> Moderate to low, if exposed, the community would be somewhat protected;</div><div><input type="checkbox"/> Very high*, if exposed, the community would be generally protected; or</div></div>



# Biological Risk Group and Containment Level Assessment

☐ Unknown.

\*Note. It is rare for the level of protection in the community to be very high. For example, community protection against Measles virus is very high because there is a highly effective vaccine and the majority of Canadians are vaccinated.

## 5. Communicability (Human and Animal Community Risk)

### Assessment of Human Communicability Indicators

Assess the indicator questions and use these to rate the likelihood of human-to-human transmission by direct or indirect contact. Use the “Rationale” section under each question to substantiate your analysis with a description and corresponding references. Note that route of infection (e.g., ingestion, inhalation) only partially addresses the likelihood of human-to-human transmission. For example, an environmental fungus may be likely to produce infection through inhalation of environmental spore, but not transmit from person-to-person, directly or indirectly. Other modes of transmission (e.g., vertical) can be noted but will not impact the final RG classification.

1. What is the likelihood of infection or disease arising from ingestion? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
2. What is the likelihood of infection or disease arising from injection (e.g., accidental or intentional inoculation, penetrating wounds)? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
3. What is the likelihood of infection or disease arising from arthropod vectors (e.g., through bites of infected arthropod species, such as mosquitoes and ticks)? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
4. What is the likelihood of infection or disease arising from contact of the agent with intact skin? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
5. What is the likelihood of infection or disease arising from contact of the agent with mucous membranes or damaged skin? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
6. What is the likelihood of infection or disease arising from inhalation of the agent (e.g., large or small droplet aerosols, spores)? <input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown  Rationale:
6. What is the likelihood of disease arising from exposure to affected animals, through either direct or indirect contact? <input type="checkbox"/> Not zoonotic <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, common mode of transmission  Rationale:
Based on the analysis of the <b>Human Communicability Indicators</b> above, rate the likelihood of human-to-human transmission by the following modes of transmission (more than one may be applicable).



## Biological Risk Group and Containment Level Assessment

Direct Contact (Casual)
<input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Direct Contact (Intimate)
<input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Indirect Contact (Fomites)
<input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Indirect Contact (Vectors)
<input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown

### Assessment of Animal Communicability Indicators

Assess the indicator questions and use these to rate the likelihood of animal-to-animal transmission by direct or indirect contact. Use the “Rationale” section under each question to substantiate your analysis with a description and corresponding references. Note that route of infection (e.g., ingestion, inhalation) only partially addresses the likelihood of animal-to-animal transmission. For example, an environmental fungus may be likely to produce infection through inhalation of environmental spore, but not transmit from animal-to-animal, directly or indirectly. Other modes of transmission (e.g., vertical) can be noted but will not impact the final RG classification.

1. What is the likelihood of infection or disease arising from ingestion?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
2. What is the likelihood of infection or disease arising from injection (e.g., accidental or intentional inoculation, penetrating wounds)?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
3. What is the likelihood of infection or disease arising from arthropod vectors (e.g., through bites of infected arthropod species, such as mosquitoes and ticks)?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
4. What is the likelihood of infection or disease arising from contact of the agent with intact skin?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
5. What is the likelihood of infection or disease arising from contact of the agent with mucous membranes or damaged skin?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
6. What is the likelihood of infection or disease arising from airborne transmission (e.g., large or small droplet aerosols, spores)?
<input type="checkbox"/> None <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, preferred route <input type="checkbox"/> Unknown
Rationale:
7. What is the likelihood of disease arising from exposure to affected humans, through either direct or indirect contact?
<input type="checkbox"/> Not zoonotic <input type="checkbox"/> Low, unlikely <input type="checkbox"/> Moderate, possible <input type="checkbox"/> High, common mode of transmission
Rationale:





## Biological Risk Group and Containment Level Assessment

Based on the analysis of the Animal Communicability Indicators above, rate the likelihood of animal-to-animal transmission by the following modes of transmission (more than one may be applicable).
Direct Contact (Casual) <input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Direct Contact (Intimate) <input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Indirect Contact (Fomites) <input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown
Indirect Contact (Vectors) <input type="checkbox"/> None <input type="checkbox"/> Unlikely <input type="checkbox"/> Possible <input type="checkbox"/> Likely <input type="checkbox"/> Unknown

### Assessment of Public Health and Economic Impact of New and/or Emerging Human Pathogens (Human Community Risk)

Complete this section **only for new or emerging human pathogens**. New or emerging pathogens, including engineered or reconstructed pathogens, may pose unique risks to the public. **Economic impact** refers to the costs associated with things like treating disease, hospitalization and long term care, and lost wages due to missed work. **Public health impact** refers to the ability of a pathogen to infect, cause disease, transmit among, and produce serious disease or death in people. Use the Rationale section under each question to substantiate your analysis with a description and corresponding references. **If you identify a new or emerging pathogen, please contact the Public Health Agency of Canada and, for emerging animal pathogens, the Canadian Food Inspection Agency to validate your risk assessment.**

1) Is the agent a new or emerging pathogen? If yes, complete the remainder of this section. If no, proceed to Section 7 (Host Range, Natural Distribution, and Economic Impact). <input type="checkbox"/> No <b>(Proceed to Section 7)</b> <input type="checkbox"/> Yes <b>(Provide detailed rational for questions 2 and 3 below)</b>  Rationale:
2) Would there be a significant impact on the economy if the pathogen were released from the laboratory (e.g., costs related to hospitalization, drugs, vaccination, and/or lost work as a result of illness)? <input type="checkbox"/> No, the anticipated economic impact would not be very high <input type="checkbox"/> Yes, <b>very high economic impact would be anticipated</b> if the pathogen were released from the laboratory  Rationale:
3) Would there be a significant impact on public health if the pathogen were released from the laboratory (e.g., significant number of cases, high health care burden)? <input type="checkbox"/> No, the anticipated public health impact would not be very high <input type="checkbox"/> Yes, <b>very high public health impact would be anticipated</b> if the pathogen were released from the laboratory  Rationale:
Based on the analysis of the New and/or Emerging Pathogen Human Pathogen Indicators above, what is the predicted impact of the release of the pathogen from a laboratory on public health or the economy: <input type="checkbox"/> Low to moderate, release from the laboratory is unlikely to have a significant impact on public health and/or the economy; or <input type="checkbox"/> Significant, release from the laboratory is likely to have a significant impact on public health and/or the economy.



Biological Risk Group and Containment Level Assessment

6. Host Range, Natural Distribution, and Economic Impact (Animal Community Risk)

Assessment of Host Range, Natural Distribution, and Economic Impact Indicators for Natural Animal Hosts

Assess the indicator questions and use these to rate the economic impact of releasing the pathogen from the laboratory on the natural animal host population. Use the Rationale section under each question to substantiate your analysis with a description and corresponding references.

1) How broad is the range of natural animal hosts that are susceptible to disease (host range)? Common classes: Amphibia, Aves, Chondrichthyes, Mammalia, Osteichthyes, Reptilia, Arachnida, Insecta.

☐ Extremely limited, single species

☐ Limited, single order

☐ Broad, single class

☐ Very broad, multiple classes

☐ Unknown

Rationale:

Are the natural host species in Canada?

☐ Natural host species are not in Canada

☐ Natural host species are present in restricted regions in Canada

☐ Natural host species are present throughout Canada

☐ Unknown

Rationale:

What is the natural distribution of the agent in Canada?

☐ Endemic in Canada

☐ Found infrequently in Canada; rare imported cases or limited natural distribution

☐ Found in Canada, but regionally restricted

☐ Not present in Canada

☐ Unknown

Rationale:

Considering animals in their order of economic importance\*, what is the combined economic value of the natural animal host(s)?

☐ None/Not Applicable

☐ Minimal

☐ Moderate

☐ Significant

☐ Unknown

Rationale:

Considering animals in their order of economic importance\*, what is the combined economic value of the other animal host(s), for example experimentally infected animals?

☐ None/Not Applicable

☐ Minimal

☐ Moderate

☐ Significant

☐ Unknown

Rationale:

Based on the analysis of the **Host range, Natural Distribution, and Economic Impact Indicators** above, the economic impact of release on the natural animal host population is:

☐ None

☐ Minimal

☐ Moderate

☐ Significant

☐ Unknown

\* The Canadian Food Inspection Agency (CFIA) has classified animals in terms of their economic value to Canada as follows:

1. Highest value livestock industries: bovine, equine, porcine, poultry, crustaceans, finfish (wild and farmed).
2. Medium value livestock industries: small ruminants (sheep and goats), bees, molluscs, other farmed

## Biological Risk Group and Containment Level Assessment

ruminants (cervids, bison).

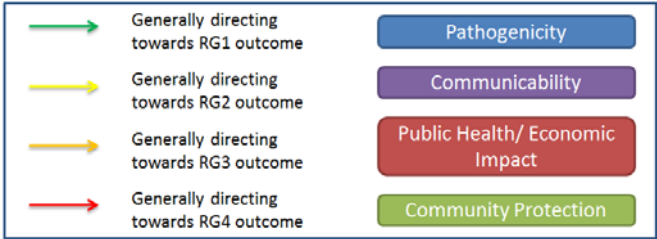
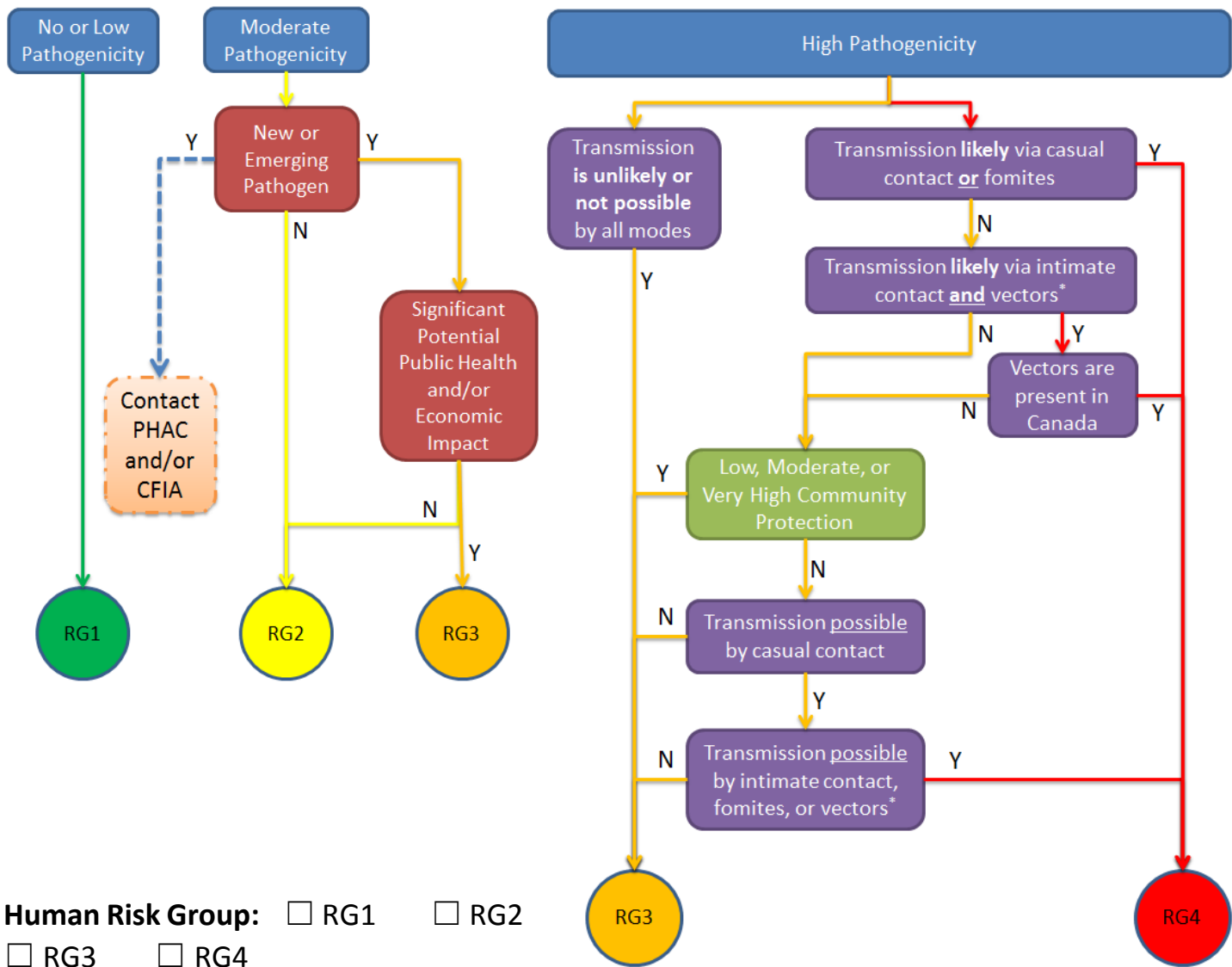
- Lowest value livestock industries and non-livestock animals: lagomorphs (rabbits), companion animals (dogs, cats, etc), reptiles, amphibians, rodents, primates.

### 7. Risk Group Decisions

The risk group reflects the risk posed to the human (Human RG) and Animal (Animal RG) populations. If the human and animal RG values differ, **the higher RG value dictates the level of containment required to work with the agent**. In almost all cases, the RG value and CL values are the same (i.e., a RG3 agent will be handled in a CL3 lab, as described in the Canadian Biosafety Standards). In rare cases, the Public Health Agency of Canada will issue Biosafety Directives that outline specific derogations of containment for certain pathogens and/or activities (<http://www.phac-aspc.gc.ca/lab-bio/res/adv-avis/index-eng.php>).

#### Human Risk Group Decision

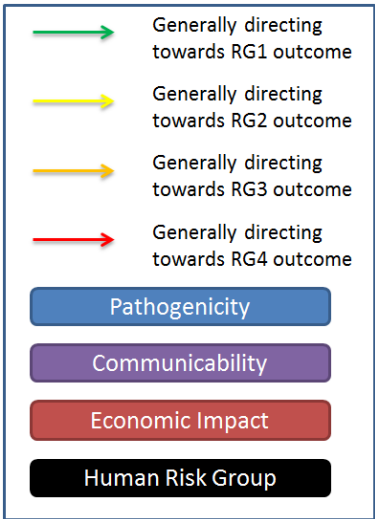
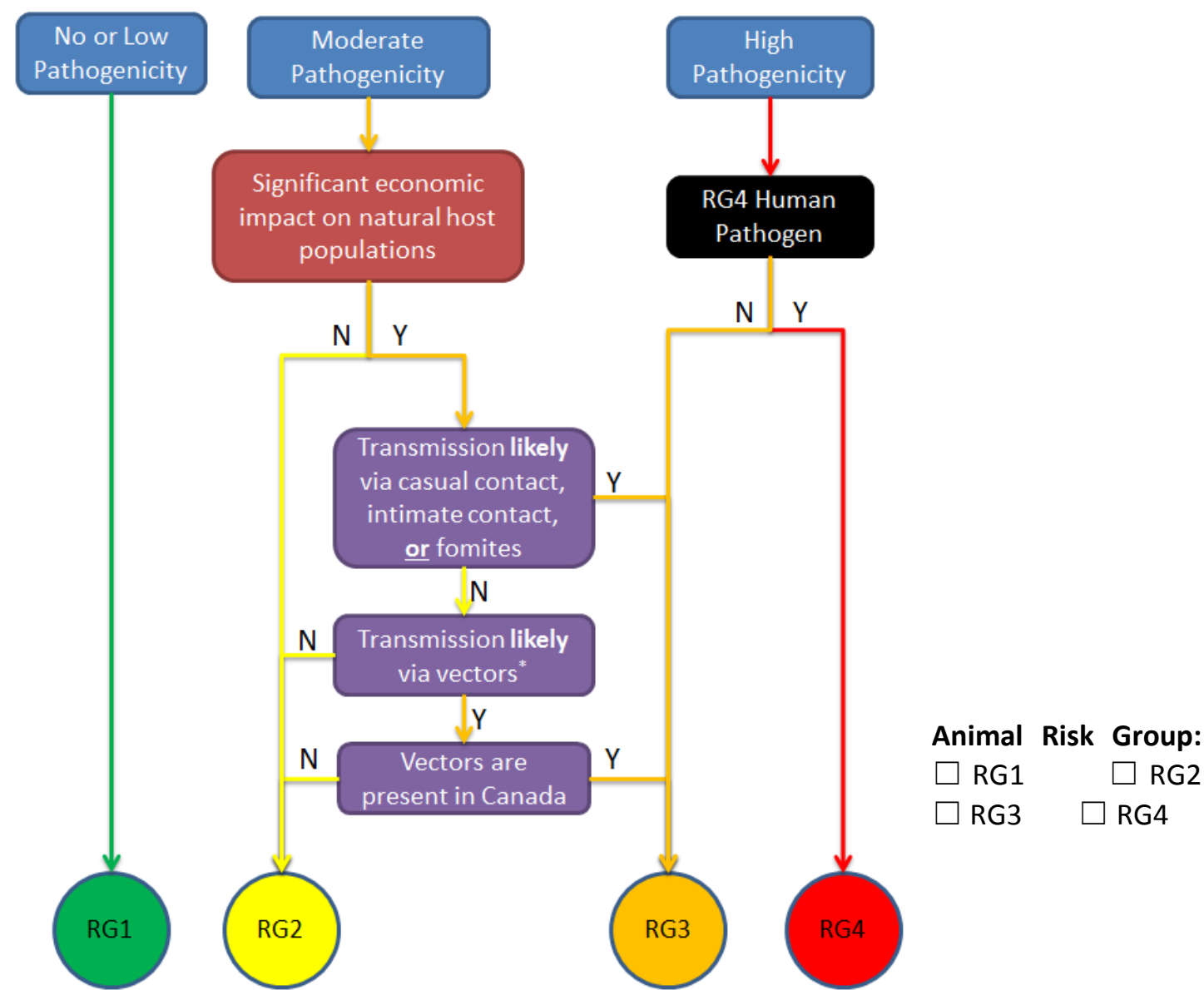
Use the decision tree to determine the risk group based on your overall rating of each of the **human** risk factor indicators.



# Biological Risk Group and Containment Level Assessment

## Animal Risk Group Decision

Use the decision tree to determine the risk group based on your overall rating of each of the **animal** risk factor indicators.



## 8. References

- All information provided in the risk assessment should be cited fully, using the highest quality data available.
- High quality data means it was sufficient for a thorough analysis of all elements of the risk assessment. High quality data sources include information from clinical trials and standardized studies.
  - Medium quality data means it was sufficient for a thorough analysis of some elements of the risk assessment but that there were some data gaps and minor assumptions were made. Medium quality data sources include peer-reviewed publish literature and edited literature.
  - Low quality data means it was insufficient for a thorough risk assessment and that there were major data gaps and major assumptions were made. Low quality data sources include expert opinion, independent communications, and uncited websites.