

Muse Safety Protocol Checklist

Directions: Use this guidance checklist to ensure that you have all required parts entered and information captured in Muse for your Protocol. Please contact IBC@uams.edu if you have any questions.

Required Muse Additions

The following headings **must** be added to either or both the **Biohazards** and **Chemical Summary** sections to ensure all safety information is captured. It is recommended to copy and paste the following headings into the text box and describe them as applicable to your research:

1. **Description:** how the hazards indicated your research will be used, what testing will be performed, etc. (Note: If not already described in another section, such as Summary of Research).
2. **Transportation of Hazards:** Primary & Secondary containment, leakproof, biohazard symbol or other appropriate hazard communication, and labelled with agent name, PI name, and PI emergency phone number (spill kit location if necessary).
3. **Centrifugation & Aerosol Prevention:** safety cups/sealed rotor use in centrifuge and vortex use.
4. **Hazard Containment & Precautions:** when will a BSC/CFH be used, PPE required for work, & splash precautions.
5. **DOT/IATA Shipping:** DOT/IATA requirements are met & appropriate containers are used. Indicate who will be performing shipping and ensure their training is documented in 'Protocol Team Members'. (Can mark N/A if not applicable).
6. **Signage:** Biohazard Signs (Agent Name, Hazards, & PPE Required), Agent Labelling, and Biohazard Door Sign (Agent Name, PI Name, PI Emergency Contact Phone Number, and Required PPE) are on the laboratory door and equipment used.
7. **Animal Use:** If applicable, describe how the agents will be used in animal (which biohazard will be used in which strain). Also indicate the **AUP** number or if an **AUP** has been submitted.
8. **Animal Facilities:** Cage Cards, Door Signs, & PPE Use for DLAM.

9. **Biosafety Cabinet/Chemical Fume Hood:** Indicate the Biosafety Cabinet (BSC) and/or Chemical Fume Hood (CFH) location and certification date in this section for all safety cabinets used.

10. **Laboratory Specific Training:** Indicate who will train the laboratory staff on specific lab procedures for all procedures in this experiment. Also indicate how the training will be documented and if there will be annual retraining.

Note: Can upload training documentation in Protocol Team Member: External Team Member Information or in Supporting Documents.

Note: BSC are not certified by UAMS. BSC are certified by an external company.

Note: Recommend using tables A- Laboratory Facilities and B- Animal Facilities from your paper protocol, if applicable.

Section Overviews

Note: Depending upon your agents, different sections will become available. Some sections below may not be applicable to your protocol and can be left blank or removed. Use the Navigation pane to move from section to section as needed.

Basic Information & Funding:

Basic Information:

<input type="checkbox"/> 1. Title of Protocol	Long descriptive title (Title of Protocol & Short Title can be the same).
<input type="checkbox"/> 2. Short Title	Title that is displayed in Muse. If applicable, include the BP# in parenthesis at the end (BP#). Note: example, 'Infection Response (BP#)'
<input type="checkbox"/> 3. Summary of Research	In plain language describe exactly what you are doing. Ensure to include the who, what, when, where and why of your experiment or clinical trial. Note: Abstract from old Paper Protocol is the same, can copy and paste.
<input type="checkbox"/> 4. Select Appropriate Safety Review	Select Biosafety. Only select Radiation Safety if you are working solely with radiation and no other hazards.
<input type="checkbox"/> 5. Principal Investigator	Type and select the Principal investigator.
<input type="checkbox"/> 6. Research Location	Select which facility your research will be performed at or performed at mostly.

Protocol Team Members:

<input type="checkbox"/> 1. Identify Team Members	Click +Add to add your team members (*by required): <input type="checkbox"/> *1. Select the <u>protocol team member</u> , search using their name (last name, first name). <input type="checkbox"/> *2. Select their role(s) in research. <input type="checkbox"/> *3. Type in <u>additional roles or titles</u> , if applicable. <input type="checkbox"/> *4. Indicate if the team member is involved in laboratory or research <u>procedures</u> . Note: *Can leave #2 blank, but then #3 is required.
<input type="checkbox"/> 2. Team Member Training	UAMS and CITI Trainings will auto populate (may need to click 'Save'). Ensure trainings

	are up-to-date and meet all of your experiment's training needs.
<input type="checkbox"/> 3. External Team Member Information	Use this to capture Arkansas Children's Trainings and Other External Training Certificates.

Funding Sources:

<input type="checkbox"/> 1. Funding Organization	Click '+Add' and select the appropriate funding organization. Also, upload funding documents or files (this section is optional: use 'UAMS' as optional funding source).
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Biosafety Summary:

Biosafety Summary:

<input type="checkbox"/> 1. Items involved in the protocol	Select all of the agents involved in the protocol. If you select 'Other', describe the item in the text box. Note: select ' Other ' if your protocol involves <u>only</u> hazardous chemicals or drugs.
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Agents, Toxins, & Microorganisms:

Tissues, Blood, or Body Fluids:

<input type="checkbox"/> 1. List agent, type, and source of all Tissues, Blood, and Body Fluids.	Click '+Add' to add Tissues, Blood, or Body Fluid Agents (*by required): <input type="checkbox"/> *1. Select the Agent . <input type="checkbox"/> *2. Select the Biosafety Level (BSL) . <input type="checkbox"/> *3. Describe the use of the Agent: describe how the agent will be used from start to finish. Where it will be collected (clinical or lab space), how many patients (if applicable; can approximate), processed (BSC, centrifuge, vortex used?), tested, shipped, etc., and how. <input type="checkbox"/> *4. Where are you obtaining the material from : example, humans, animals, patients, manufacturer, etc. <input type="checkbox"/> *5. Select the storage locations . Note: use the % to search for your room number. Example: %1234 <input type="checkbox"/> *6. Select the usage locations . <input type="checkbox"/> 7. Indicate the supplier (if applicable): who the manufacturer is.
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	<ul style="list-style-type: none"> <input type="checkbox"/> 8. Indicate maximum quantity stored in the lab space, if known. <input type="checkbox"/> 9. Indicate personnel that will be handling the material, if applicable: example, this field is necessary if only certain members of the research team are trained or have the qualifications to handle the agent. <input type="checkbox"/> 10. Experimental concentration, if applicable. <input type="checkbox"/> *11. Is agent used in animals? <input type="checkbox"/> *12. Is agent used in humans? <input type="checkbox"/> *13. Is agent recombinant or synthetic?
<ul style="list-style-type: none"> <input type="checkbox"/> 2. Describe any tissues transplanted between species, if applicable. 	<p>Indicate if you are transplanting tissues or mark N/A.</p>
<ul style="list-style-type: none"> <input type="checkbox"/> 3. Describe the quantity of tissues and volumes of fluids to be used. 	<p>Indicate the quantities of tissues and volumes to be used throughout your experiment, in detail: example, 3 mL of blood will be collected from approximately 50 patients for a maximum quantity of 150 mL.</p>

Primary Cells or Cell Lines:

<ul style="list-style-type: none"> <input type="checkbox"/> 1. List agent, type, and source of all Primary Cells or Cell Lines. 	<p>Click '+Add' to add Primary Cells or Cell Lines (*by required):</p> <ul style="list-style-type: none"> <input type="checkbox"/> *1. Select the Agent. <input type="checkbox"/> *2. Select the Biosafety Level (BSL). <input type="checkbox"/> *3. Describe the use of the Agent: describe how the agent will be used from start to finish. Where it will be collected (clinical or lab space), how many patients (if applicable; can approximate), processed (BSC, centrifuge, vortex used?), experimental use, etc., and how. <input type="checkbox"/> *4. Where are you obtaining the material from: example, humans, animals, patients, manufacturer, etc. <input type="checkbox"/> *5. Select the storage locations. Note: use the % to search for your room number. Example: %1234 <input type="checkbox"/> *6. Select the usage locations. <input type="checkbox"/> 7. Indicate the supplier (if applicable): who the manufacturer is.
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	<input type="checkbox"/> 8. Indicate maximum quantity stored in the lab space, if known. <input type="checkbox"/> 9. Indicate personnel that will be handling the material, if applicable: example, this field is necessary if only certain members of the research team are trained or have the qualifications to handle the agent. <input type="checkbox"/> 10. Experimental concentration , if applicable. <input type="checkbox"/> *11. Is agent used in animals? <input type="checkbox"/> *12. Is agent used in humans? <input type="checkbox"/> *13. Is agent recombinant or synthetic?
<input type="checkbox"/> 2. List other mammalian species cell lines.	Indicate which, if any, of your cell lines listed above are non-human mammalian cells. Also list any cell line transduced derivatives.
<input type="checkbox"/> 3. List other non-mammalian species cell lines.	Indicate which, if any, of your cell lines listed above are non-mammalian cells.
<input type="checkbox"/> 4. Identify cultures in volumes over 10 liters.	Indicate if you have any cultures more than 10 liters or mark N/A.

Bacteria, Yeasts, Fungi, or Parasites:

<input type="checkbox"/> 1. List agent, type, and source of all Microorganisms: Bacteria, Yeasts, Fungi, or Parasites.	Click '+Add' to add Tissues, Blood, or Body Fluid Agents (*by required): <input type="checkbox"/> *1. Select the Agent . <input type="checkbox"/> *2. Select the Biosafety Level (BSL) . <input type="checkbox"/> *3. Describe the use of the Agent: describe how the agent will be used from start to finish. <input type="checkbox"/> 4. Indicate the strain , if applicable. <input type="checkbox"/> *5. Where are you obtaining the material from : example, humans, animals, patients, manufacturer, etc. <input type="checkbox"/> *6. Select the storage locations . Note: use the % to search for your room number. Example: %1234 <input type="checkbox"/> *7. Select the usage locations . <input type="checkbox"/> 8. Indicate the supplier (if applicable): who the manufacturer is.
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	<ul style="list-style-type: none"> <input type="checkbox"/> 9. Indicate maximum quantity stored in the lab space, if known. <input type="checkbox"/> 10. Indicate personnel that will be handling the material, if applicable: example, this field is necessary if only certain members of the research team are trained or have the qualifications to handle the agent. <input type="checkbox"/> 11. Experimental concentration, if applicable. <input type="checkbox"/> *12. Is agent used in animals? <input type="checkbox"/> *13. Is agent used in humans? <input type="checkbox"/> *14. Is agent recombinant or synthetic?
<input type="checkbox"/> 2. Describe other microorganisms	<p>Optional: describe other microorganisms being used that are not primary parts of the experiment or mark N/A. Note: Can include other microorganisms in the above section.</p>

Viruses or Prions:

<input type="checkbox"/> 1. List agent, type, and source of all Viruses or Prions.	<p>Click +Add to add Tissues, Blood, or Body Fluid Agents (*by required):</p> <ul style="list-style-type: none"> <input type="checkbox"/> *1. Select the Agent. <input type="checkbox"/> *2. Select the Biosafety Level (BSL). <input type="checkbox"/> *3. Describe the use of the Agent: describe how the agent will be used from start to finish. <input type="checkbox"/> 4. Indicate the strain, if applicable. <input type="checkbox"/> *5. Where are you obtaining the material from: example, humans, animals, patients, manufacturer, etc. <input type="checkbox"/> *6. Select the storage locations. Note: use the % to search for your room number. Example: %1234 <input type="checkbox"/> *7. Select the usage locations. <input type="checkbox"/> 8. Indicate the supplier (if applicable): who the manufacturer is. <input type="checkbox"/> 9. Indicate maximum quantity stored in the lab space, if known. <input type="checkbox"/> 10. Indicate personnel that will be handling the material, if applicable: example, this field is necessary if only certain members of the research team are trained
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	<p>or have the qualifications to handle the agent.</p> <p><input type="checkbox"/> 11. Experimental concentration, if applicable.</p> <p><input type="checkbox"/> *12. Is agent used in animals?</p> <p><input type="checkbox"/> *13. Is agent used in humans?</p> <p><input type="checkbox"/> *14. Is agent recombinant or synthetic?</p>
<p><input type="checkbox"/> 2. Describe other Viruses or Prions</p>	<p>Optional: describe other viruses or prions being used that are not primary parts of the experiment or mark N/A.</p> <p>Note: Can include other viruses or prions in the above section.</p>

Select Agents & Toxins or Toxins:

<p><input type="checkbox"/> 1. List agent, type, and source of all Select Agents & Toxins or Toxins.</p>	<p>Click '+Add' to add Tissues, Blood, or Body Fluid Agents (*by required):</p> <p><input type="checkbox"/> *1. Select the Agent.</p> <p><input type="checkbox"/> *2. Select the Biosafety Level (BSL).</p> <p><input type="checkbox"/> *3. Describe the use of the Agent: describe how the agent will be used from start to finish.</p> <p><input type="checkbox"/> *4. Where are you obtaining the material from: example, humans, animals, patients, manufacturer, etc.</p> <p><input type="checkbox"/> *5. Select the storage locations. Note: use the % to search for your room number. Example: %1234</p> <p><input type="checkbox"/> *6. Select the usage locations.</p> <p><input type="checkbox"/> 7. Indicate the supplier (if applicable): who the manufacturer is.</p> <p><input type="checkbox"/> 8. Indicate maximum quantity, if known.</p> <p><input type="checkbox"/> 9. Indicate personnel that will be handling the material, if applicable: example, this field is necessary if only certain members of the research team are trained or have the qualifications to handle the agent.</p> <p><input type="checkbox"/> 10. Experimental concentration, if applicable.</p> <p><input type="checkbox"/> *11. Is agent used in animals?</p> <p><input type="checkbox"/> *12. Is agent used in humans?</p>
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	<input type="checkbox"/> *13. Is agent recombinant or synthetic?
<input type="checkbox"/> 2. Does this protocol involve an excluded select agent or toxin?	Indicate yes or no as appropriate.
<input type="checkbox"/> 3. Does this protocol involve any select agents or toxins in exempted amounts?	Indicate yes or no as appropriate.
<input type="checkbox"/> 4. Describe the volumes of cultures to be used for any select agents or toxins.	If needed, indicate in detail the quantities of Select Agents & Toxins or Toxins or mark N/A.

Biohazards:

<input type="checkbox"/> 1. Summary of each agent, toxin, or microorganism that will be used in this protocol.	Verify that <u>all biohazard agents</u> that will be used in your protocol are listed.
<input type="checkbox"/> 2. Provide a description of any agents, toxins, or microorganisms indicated above.	Indicate any procedural or descriptive information for the agents that has not already been specified, such as: <ul style="list-style-type: none"> <input type="checkbox"/> Description: how the hazards indicated your research will be used, what testing will be performed, etc. (Note: If not already described in another section, such as Summary of Research). <input type="checkbox"/> Transportation of Hazards: Primary & Secondary containment, leakproof, biohazard symbol or other appropriate hazard communication, and labelled with agent name, PI name, and PI emergency phone number. <input type="checkbox"/> Centrifugation & Aerosol Prevention: safety cups/sealed rotor use in centrifuge and vortex use.

	<ul style="list-style-type: none"> <input type="checkbox"/> Hazard Containment & Precautions: BSC use, & splash precautions. <input type="checkbox"/> DOT/IATA Shipping: DOT/IATA requirements are met & appropriate containers are used. Indicate who will be performing shipping and ensure their training is documented in 'Protocol Team Members'. <input type="checkbox"/> Signage: Biohazard Signs (Agent Name, Hazards, & PPE Required), Agent Labelling, and Biohazard Door Sign (Agent Name, PI Name, PI Emergency Contact Phone Number, and Required PPE). <input type="checkbox"/> Animal Use: Describe how the agents will be used in animal (which biohazard will be used in which strain). Note: Can describe animal use in #3 for each agent. <input type="checkbox"/> Animal Facilities: Cage Cards, Door Signs, & PPE Use for DLAM. <input type="checkbox"/> *BSC: If applicable, indicate the Biosafety Cabinet (BSC) <u>location</u> and <u>certification date</u> in this section for all BSC used. Note: BSC are not certified by UAMS. BSC are certified by an external company. <input type="checkbox"/> *Laboratory Specific Training: If applicable, indicate who will train the laboratory staff on specific lab procedures for the <u>biohazards</u> in this experiment. How the training will be documented and if there will be annual retraining. Note: Can upload training documentation in Protocol Team Member: <u>External Team Member Information</u> or in <u>Supporting Documents</u>.
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	Note: Recommend using tables B, C, & D from Section V: Research Elements from your paper protocol, if applicable.
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Recombinant & Synthetic Nucleic Acids:

Recombinant or Synthetic Nucleic Acids Usage:

<input type="checkbox"/> 1. Does research with recombinant or synthetic nucleic acids involve the use of.	Select the appropriate NIH Guideline for your research involving recombinant or synthetic nucleic acids (r/sNA). Note: see the NIH Guidelines for more information or if you have a past paper protocol, section A. iii.
<input type="checkbox"/> 2. If none of these apply, describe.	If you did not select an appropriate NIH Guideline above, indicate why and describe your research. Note: Write N/A or leave blank if this text box is not applicable.

Recombinant or Synthetic Nucleic Acid Work Description:

<input type="checkbox"/> 1. Describe any work involving recombinant or synthetic nucleic acid molecules. Include information about host-vector systems, genes, and procedures.	Give an overview of <u>what</u> you are doing with recombinant or synthetic nucleic acids (r/sNA) and the risk attenuation . For example: Cloning, PCR, expression in a microbe, use in a tissue culture, or use in an organism.
<input type="checkbox"/> 2. For each experiment, list genes, inserts, gene products, and key regulatory elements to be cloned.	List the following types of items to be cloned in your research in the text box: <input type="checkbox"/> Genes (explain acronyms) <input type="checkbox"/> Inserts <input type="checkbox"/> Gene Products <input type="checkbox"/> Key Regulatory Elements
<input type="checkbox"/> 3. Provide a brief description of gene activity and indicate the species of origin for each.	Indicate the following for all genes listed above: <input type="checkbox"/> Description of Gene Activity: state what each Gene listed above does. Note: Give the Nature of Insert or Protein Expressed (Toxin, antibiotic resistance/selection marker, virulence factor, reporter, oncogene, transcription factor, etc.)

	<input type="checkbox"/> Species of Origin (Genus and species, strain) for <u>each</u> Gene Activity listed. Note: Use the table in Section A.i. of your past paper protocol, column 3 and column 1, if applicable.
<input type="checkbox"/> 4. Describe all vectors (plasmids, viruses, RNA/DNA constructs) to be used	List out all vectors to be used and describe what they will be used for: Gene Transfer Method, Vector & Biosafety Level, Vector Function (cloning, protein expression, etc.), & Risk Attenuation. Note: Use the table in Section A.ii. of your past paper protocol, if applicable.
<input type="checkbox"/> 5. Upload vector map.	Upload your vector map. Note 1: If you are using/generating multiple r/sNA, number them and use the corresponding numbers throughout the 'Recombinant Nucleic Acid Work Description' section for the corresponding vectors, hosts, vector maps, etc. Note 2: For multiple vector maps, copy and paste them into a Word document, print to PDF, and then upload the PDF file.
<input type="checkbox"/> 6. For each experiment identify all applicable host systems to be used.	List & explain the use of each of the applicable items: <ul style="list-style-type: none"> <input type="checkbox"/> Packaging Systems (Tables A.i-iii from paper protocol) <input type="checkbox"/> Microbes (Table B from paper protocol) <input type="checkbox"/> Tissue/Cell Culture (Table C from paper protocol) <input type="checkbox"/> Plant/Animal (Table D from paper protocol)
<input type="checkbox"/> 7. Describe any attempts to express foreign genes from living organisms.	Indicate in detail foreign genes to be expressed & list what organism they are from.
<input type="checkbox"/> 8. List any proteins produced.	List proteins.

Note: If you have a previous paper protocol, ensure all information from Section A is captured in the Recombinant Nucleic Acid section of Muse.

Human Gene Transfer/Human Clinical Trials:

<input type="checkbox"/> 1. Provide a brief description of the	List or indicate the goals of your research (why are you doing this study).
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overall goals and research activities.	
<input type="checkbox"/> 2. Provide a summary of biosafety-relevant preclinical and clinical studies associated with this protocol <i>*Attach Summary Documents*</i>	<p>Describe any biosafety related information for your study. Click the ? button for help information (attach supporting documents and lab procedures as needed).</p> <p>Note: if this protocol is only for prep and shipping of human specimens, indicate that the collection will be performed in the clinical area and that lab staff are following and have been trained in safe bloodborne pathogen handling procedure.</p>
<input type="checkbox"/> 3. Provide a description of the materials.	<p>List any materials used and describe their use. If only shipping human body fluids or tissue, indicate that.</p>
<input type="checkbox"/> 4. Describe specific procedures for clinical monitoring of the biohazard used in the protocol (e.g. potential for shedding)	<p>List procedures for staff to take if exposed to biohazards from the human trial.</p> <p>Note: if this protocol is only for prep and shipping of human specimens, this section may be similar or match #1 in the 'Exposure Assessment and Protective Equipment' section.</p>
<input type="checkbox"/> 5. Attach all relevant standard operating procedures (SOPs).	<p>Upload any other SOPs or brochures for this clinical trial (can upload multiple files).</p> <p>Note: Can upload PCRU SOP here or in 'Supporting Documents'.</p>
<input type="checkbox"/> 6. Attach clinical protocols.	<p>Upload the clinical protocols or the study details here (can only upload one file).</p>
<input type="checkbox"/> 7. Additional Documents.	<p>Any additional documentation, IRB approval letter, etc. can go here.</p>
<input type="checkbox"/> 8. Has IRB review been initiated in CLARA?	<p>Indicate yes or no as appropriate.</p> <p>Note: Ensure to complete the Manage Related IRB Studies to input information about your Human Clinical Trial, located on your protocol's homepage.</p>
<input type="checkbox"/> 9. CLARA Number.	<p>Type in your CLARA number (can list them)</p>

Animals and Genetically Modified Animals:

Animals:

<input type="checkbox"/> 1.a. Identify the species to be used (click the three dots).	<p>Select each of the animal species to be used.</p>
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	Note: email Biosafety Officers if you need to add a species.
<input type="checkbox"/> 1.b. Identify the locations where animals are being housed or used (click the three dots).	Select the location(s) that the animals listed above will be house and used. Note: use the % to search for your room number. Example: %1234
<input type="checkbox"/> 2. Are the animals used in the experiment immunocompromised?	Indicate yes or no as appropriate.
<input type="checkbox"/> 3. If yes, describe how.	List all animal strains to be used and indicate which ones are immunocompromised. Describe how each animal strain will be used. Indicate AUP number(s) , if known or if an AUP has been submitted.
<input type="checkbox"/> 4. Which of the following present exposure risks to the protocol team members or animal care personnel?	Select each that applies for your animals (consider the biohazards) for any animal handling staff, such as DLAM or your laboratory team.

Genetically Modified Animals: DNA Source

<input type="checkbox"/> 1. Describe the transgene and remaining vector sequences.	List each transgenic animal, their transgenes and remaining vector sequences. Describe the use or purpose of each.
<input type="checkbox"/> 2. Will you be purchasing, breeding, or obtaining transgenic rodents from an external laboratory or have you done so in the past?	Have you or will you purchase or receive transgenic rodents for this protocol? Indicate yes or no as applicable.
<input type="checkbox"/> 3. Will this protocol involve creating a transgenic strain on site at a BSL-1 containment level?	Are you generating BSL-1 transgenic rodents in your lab space in this protocol? Indicate yes or no as applicable.

<input type="checkbox"/> 4. What is the source of the DNA?	Select all that apply.
<input type="checkbox"/> 5. Are human or animal pathogens to be used as a host-vector system?	Indicate yes or no as applicable.
<input type="checkbox"/> 6. Does this experiment use viruses?	Indicate yes or no as applicable.

Gene Transfer: Transgenic Strain

Note: Only complete this section if **you are creating transgenic animals** in your laboratory. Do **not** complete this section if you are purchasing or receiving the transgenic animal strain(s) from a supplier or another laboratory.

<input type="checkbox"/> 1. Describe why it is necessary to generate transgenics to conduct this research	Relevant to this protocol, why are you creating a transgenic animal strain.
<input type="checkbox"/> 2. Describe the goals of the research and all procedures used.	Briefly describe how the transgenic animal strains generated help achieve your research goals. List all procedures and all biohazards to be used with the transgenic animals.
<input type="checkbox"/> 3. Describe how this research relates to the NIH Guidelines for Research Involving rDNA.	Describe the animal genome alteration and the Biosafety Level (BSL), does this fall under Section III-E-3 or III-D-4. Note: Click the NIH Guidelines for more information and help (Read Section III-E-3: Experiments Involving Transgenic Rodents).
<input type="checkbox"/> 4. Does this protocol involve the alteration of the germ line of the animal?	Indicate yes or no as appropriate.
<input type="checkbox"/> 5. How will the DNA be introduced?	Describe how the DNA will be introduced to create the transgenic animal, such as DNA microinjection, embryonic stem cell-mediated gene transfer, or retrovirus-mediated gene transfer.

Plants:

Transgenic Plants:

<input type="checkbox"/> 1. Does your protocol involve the use of transgenic plants	Indicate yes or no as appropriate.
<input type="checkbox"/> 2. If yes, provide a description of each transgenic plant species.	Describe each transgenic plant species used: how you will use in your protocol, transport, containment (BSC), disinfect, store, quantity, biosafety level, etc.

Risk Management:

Risk Group and Containment Practices:

<input type="checkbox"/> 1. What is the highest risk group level of the biological agents and materials you will use in the proposed research?	Classify your protocol as risk group 1-4. Note: Most UAMS/ACRI protocols are Risk Group 2 (RG2). Click the blue link to use the NIH Guidance for help.
<input type="checkbox"/> 2. What are the highest biosafety containment practices required for the research activities covered by this protocol?	For your protocol, select <u>only</u> the biosafety levels that apply to the agents you are using (rDNA, plants, animals, etc.).

Exposure Assessment and Protective Equipment:

<input type="checkbox"/> 1. Describe consequences of exposure or release of agents used to humans, animals, and plants.	For each agent listed in your protocol describe what to do if you have a skin, mucous membrane, eye, inhalation, or other exposure: <ul style="list-style-type: none"> <input type="checkbox"/> Biohazard Agents: for all listed in Agents, Toxins, & Microorganisms section <input type="checkbox"/> r/sNA or rDNA: for all listed in Recombinant & Synthetic Nucleic Acids section <input type="checkbox"/> Animals: for all listed in the Animals section Note: Ensure that all potential exposures, such as animal bedding, checked in #4 of the Animals section are addressed. <input type="checkbox"/> Plants: for all listed in the Plants section
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	<p><input type="checkbox"/> Chemicals/Drugs: for all listed in the Custom Pages</p> <p>Note: use or cite the UAMS Biosafety Manual Can use excerpt in your UAMS protocol:</p> <p>Skin, Mucous Membrane, or Injury Exposure: All exposures to recombinant or synthetic nucleic acid molecules or recombinant pathogens must be reported to the laboratory supervisor or PI immediately, as well as the Biological Safety Officer (501-686-5536). If the exposure results in an emergency, call the UAMS Call Center at 501-526-0000 or 911 to request help. Otherwise, the following steps should be taken: If there is an agent-specific protocol for exposure, such as those for HIV or Hepatitis B, follow those protocols.</p> <p>For skin contact exposure or injury with a contaminated instrument:</p> <ul style="list-style-type: none"> • Thoroughly wash the area with soap and water. Do not squeeze the wound to induce bleeding. • Avoid using abrasive chemical soaps or disinfectant washes, as they can cause skin abrasions and a possible additional route of entry for the agent. • Cover the wound with sterile dressing. • For mucous membranes (e.g., eyes, mouth), flush with water for a minimum of 15 minutes. • Contact UAMS Student and Employee Health as soon as possible after exposure • Exposed personnel must complete the Employee/Student Injury and Incident Report (I&I) Form. • Employees of UAMS must call the Company Nurse Injury Hotline at 1- 855-339-1893 as soon as possible after injury or exposure. • During normal business hours, exposed individuals should contact the Preventive Occupational Environmental Medicine Clinic at 501-686-6565 (open 8 a.m. to 4:30 p.m.) • After hours, exposed individuals should contact the UAMS Emergency Department at 501-686-2085 for evaluation. <p>Any exposures to recombinant DNA must be reported to the following:</p> <ul style="list-style-type: none"> • Principal Investigator or Laboratory Director—Immediately • Biological Safety Officer (501-686-5536)—Immediately
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	<ul style="list-style-type: none"> • Institutional Biosafety Committee (IBC)— May be reported through Biological Safety Officer • NIH/OBA—Report will be coordinated by the IBC
<input type="checkbox"/> 2. Indicate the personal protective equipment that will be used.	Select all PPE that will be used at any point in your protocol.
<input type="checkbox"/> 3. If other, specify.	<input type="checkbox"/> Indicate Other PPE . Note: Also use this space to indicate if certain PPE selected above will only be used during certain procedures , such as eye protection will only be used when procedures with a splash risk are performed. Note: Also use this space to indicate what type of respirator will be used, such as N-95, PAPR, CAPR, etc.

Dual Use Research of Concern:

<input type="checkbox"/> 1. Dual use experiment categories used in this research	Select all that apply. Note: most UAMS/ACRI protocols are 'None of the above'.
<input type="checkbox"/> 2. Explain why you believe this protocol is or is not dual use research of concern	Explain your above selection. For ' None of the above ' the following statement can be used, if it is applicable to your protocol: <i>No agents are being used that are subject to the US Government DURC policies and no laboratory procedure performed will increase the harmful consequences of any agent or toxin present.</i>

Waste Management:

<input type="checkbox"/> 1. Describe the process for decontaminating biological waste.	For each agent listed in your protocol describe how the waste generated will be disposed of for all that apply: <input type="checkbox"/> Surfaces & Equipment used in your research.
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	<ul style="list-style-type: none"> <input type="checkbox"/> Biohazard Agents: for all listed in Agents, Toxins, & Microorganisms section. <input type="checkbox"/> r/sNA or rDNA: for all listed in Recombinant & Synthetic Nucleic Acids section. <input type="checkbox"/> Animal Wastes & Animal Bedding: for all listed in the Animals section. Note: Ensure that all potential wastes, such as animal bedding, is considered. <input type="checkbox"/> Plants: for all listed in the Plants section. <input type="checkbox"/> Chemicals/Drugs: for all listed in the Custom Pages. <input type="checkbox"/> *Disinfectant and Contact Time: must indicate for all disinfectants used. Example, 10% bleach for a minimum of 20 minutes (recommend wiping with 70% ethanol after to prevent corrosion). <input type="checkbox"/> *Autoclave time & temperature: if applicable. Example, 121°C for 15 minutes. <input type="checkbox"/> *Sharps: disposed of in appropriate Sharps containers. <p>Note 1: use or cite the UAMS Biosafety Manual Decontamination (Section 11) & Waste Management (Section 13). Also consider liquid vs. solid biohazard waste. Can use excerpt in your protocol:</p> <p>13.4.1 Sharps Waste All sharps must be placed into properly labeled sharps containers or other rigid, puncture-proof containers. Sharps containers must not be filled more than 2/3 full. Make sure the container is sealed, labeled, and intact. When filled, request pickup of sharps containers by contacting the UAMS Biohazard Waste Pickup Team (501-526-0000) or requesting a pickup online: http://www.uams.edu/campusop/depts/ohs/forms/biowaste.aspx.</p> <p>13.4.2 Solid Biohazardous Waste Solid waste includes cloth, plastic, and paper items that have been exposed to infectious agents hazardous to humans, animals, or plants, and solidified agarose gels. These contaminated items should be collected in appropriate biohazard waste</p>
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	<p>autoclave bags. When $\frac{3}{4}$ full, loosely tie the bag closed and secure the lid on the container. Contact the UAMS Biohazard Waste Pickup Team (5001-526-0000) to request pickup or request a pickup online http://www.uams.edu/campusop/depts/ohs/forms/biowaste.aspx.</p> <p>13.4.3 Liquid Biohazardous Waste Liquid biological waste should be collected in leak proof, rigid, durable containers for autoclaving or chemical disinfection. Autoclaved or chemically disinfected liquid wastes can be disposed of via the laboratory sink. Do not pour melted agarose down the drain. Allow it to cool and solidify, then dispose of it as solid waste in biohazard waste bags.</p> <p>Example: All Liquid waste (Cell lines, rDNA, and infected material: treated for 20 minutes with 10% bleach. All surfaces: wiped with 10% bleach and then with 70% ethanol and allowed to air dry. All solid waste: placed in red biohazard bags to be picked up by the Biohazard Waste team for offsite disposal. Reusable plastic & glassware: autoclaved at 121°C and 15 psi for 30 minutes. Chemotherapy Waste: all chemotherapy agents and chemotherapy contaminated wastes are disposed of in yellow bags and yellow containers for off-site disposal. Sharps: all sharps are placed in red Sharps containers that are sealed when full and picked up by the Biohazard Waste team to be disposed of offsite. Animal bedding & waste: Cages are dumped using a dump station into red biohazard waste and incinerated offsite. All cages are autoclaved per DLAM policy to ensure sufficient sterilization.</p> <p>Note 2: do not autoclave anything that has been treated with bleach. Note 3: radioactive or pharmaceutical chemicals used must be disposed of properly and indicated in this section.</p>
<p><input type="checkbox"/> 2. Autoclave location.</p>	<p>Indicate where your autoclave is, if use is indicated in above section.</p> <p>Note: use the above text box to indicate the location of other autoclaves if more than one is used.</p>

<p><input type="checkbox"/> 3. Describe the plans for decontamination in the event of a biological accident.</p>	<p>For each agent listed in your protocol describe how large spills, small spills, and other biological incidents will be addressed for all that apply:</p> <ul style="list-style-type: none"> <input type="checkbox"/> Biohazard Agents: for all listed in Agents, Toxins, & Microorganisms section <input type="checkbox"/> r/sNA or rDNA: for all listed in Recombinant & Synthetic Nucleic Acids section <input type="checkbox"/> Animals: for all listed in the Animals section Note: Ensure that all potential wastes, such as animal bedding, is considered. <input type="checkbox"/> Plants: for all listed in the Plants section <input type="checkbox"/> Chemicals/Drugs: for all listed in the Custom Pages <input type="checkbox"/> *Disinfectant and Contact Time: must indicate for all disinfectants used. Example, 10% bleach for a minimum of 20 minutes (recommend wiping with 70% ethanol after to prevent corrosion). <input type="checkbox"/> *Autoclave time & temperature: if applicable. Example, 121°C for 15 minutes. <input type="checkbox"/> *Sharps: disposed of in appropriate Sharps containers. <p>Note 1: use or cite the UAMS Biosafety Manual Spill Procedure (Section 10). Also consider liquid vs. solid biohazard waste.</p>
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Custom Pages:

Chemical Hygiene:

<p><input type="checkbox"/> 1. Provide relevant hygiene information for each chemical used in the protocol.</p>	<p>Click '+Add' to add Chemicals/Drugs (*by required):</p> <ul style="list-style-type: none"> <input type="checkbox"/> *1. Select the Agent. Note: select 'Not Applicable' if your protocol does not have chemicals and no other fields are required. <input type="checkbox"/> *2. Describe the <u>physical properties</u> of the Agent: describe how the agent will be used from start to finish.
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	<ul style="list-style-type: none"> <input type="checkbox"/> 3. Select the classification, if applicable. <input type="checkbox"/> *4. Describe the toxicology of the chemical being used (use SDS). Note: Must indicate if chemical is a carcinogen, mutagen, teratogen, toxin, radioactive, etc. (see paper protocol section V. table E) <input type="checkbox"/> *5. Indicate routes of exposure for this chemical or drug: ensure <u>all</u> routes of exposure are addressed in the 'Exposure Assessment and Protective Equipment' section. <input type="checkbox"/> 6. Other precautions: such as, what PPE to use and if a CFH will be used. <input type="checkbox"/> 7. Exposure Limits: indicate the Permissible Exposure Limit (PEL) from SDS. <input type="checkbox"/> 8. Monitoring requirements: Consider radiative chemicals (will wipe tests or badges be required).
<ul style="list-style-type: none"> <input type="checkbox"/> 2. Justify the use of the selected chemicals. 	<p>Indicate why you are using these chemicals/drugs.</p>

Chemical Summary:

<ul style="list-style-type: none"> <input type="checkbox"/> 1. Identify the chemicals to be used in the protocol. 	<p>Click ' Update' to add the locations for your Chemicals/Drugs (*by required):</p> <ul style="list-style-type: none"> <input type="checkbox"/> *1. Should already be prefilled by the Muse system (if 'Not Applicable' indicated, continue to #4) <input type="checkbox"/> *2. Indicate the maximum Quantity that will be used in your experiments (Max amount administered). <input type="checkbox"/> 3. Indicate the number of procedures that the chemical is involved in. Note: Leave blank if used in all procedures. <input type="checkbox"/> *4. Describe the storage location of the chemical being used. Note: select 'Not Applicable' if your protocol does not have chemicals and no other fields are required. <input type="checkbox"/> *5. Indicate the maximum Quantity that will be kept onhand.
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	<p><input type="checkbox"/> *6. Indicate the Chemical Fume Hood (CFH) certification date. Note: If the location of the CFH is different from the location in #4 or there is more than one CFH used, indicate this information in Section 2 below.</p> <p><input type="checkbox"/> *7. Lab phone number: put the phone number to call for emergencies.</p> <p><input type="checkbox"/> *8. Attachments: upload the chemical's SDS or toxicology information, if applicable.</p> <p>Note: recommend using the 'Highly Toxic Chemicals...' (table E) from your paper protocol if applicable.</p>
<p><input type="checkbox"/> 2. Describe the experiment, including procedures used in the protocol.</p>	<p>Indicate any procedural or descriptive information for the agents:</p> <ul style="list-style-type: none"> <input type="checkbox"/> Description: what each agent is or what it will be used for in your research. <input type="checkbox"/> Transportation of Hazards: Primary & Secondary containment (will a spill kit be used). <input type="checkbox"/> Centrifuge, Vortex, & Equipment Use: safety cups/sealed rotor, etc. <input type="checkbox"/> Hazard containment & precautions: CFH use, & splash precautions (BSC use, if not already indicated). <input type="checkbox"/> DOT/IATA Shipping: DOT/IATA requirements are met & appropriate containers are used. Indicate who will be performing shipping and ensure their training is documented in 'Protocol Team Members'. <input type="checkbox"/> Signage: Hazard Signs (Agent Name, Hazards, & PPE Required) & Agent Labelling. <input type="checkbox"/> Animal Use: Describe how the agents will be used in animal

	<p>(which chemical will be used in which strain).</p> <p>Note: Can describe animal use in #3 for each agent.</p> <p><input type="checkbox"/> Animal Facilities: Cage Cards, Door Signs, & PPE Use for DLAM.</p> <p><input type="checkbox"/> *Chemical Fume Hood: If applicable, indicate the Chemical Fume Hood (CFH) <u>location</u> and <u>certification date</u> in this section for all CFH used (if not captured in section #1 above).</p> <p>Note: CFH are certified by UAMS, call the call center (501-526-0000) if yours need certification.</p> <p><input type="checkbox"/> *Laboratory Specific Training: If applicable, indicate who will train the laboratory staff on specific lab procedures for the <u>chemicals</u> in this experiment. How the training will be documented and if there will be annual retraining.</p> <p>Note: Can upload training documentation in Protocol Team Member: <u>External Team Member Information</u> or in <u>Supporting Documents</u>.</p>
<p><input type="checkbox"/> 3. Describe the methods used.</p>	<p>Indicate the exact procedural steps that will be used for each method. Recommend numbering methods.</p> <p>Example: Method 1: subcutaneous mouse injection. 0.5 mg/kg Chemical will be prepared in the CFH using PPE indicated. The chemical will be transported to DLAM in a labelled secondary container per UAMS Biosafety Manual protocol. The chemical will be administered to mice in the DLAM BSC via flank subcutaneous injection. Appropriate Door Signs and Cage Cards will be placed that indicate the chemical name, hazards, PI, and PI emergency contact phone number.</p> <p>Note: recommend using the 'Highly Toxic Chemicals...' (table E) from your paper protocol if applicable.</p>
<p><input type="checkbox"/> 4. Will liquid N₂ or cryogenic liquid be used?</p>	<p>Indicate yes or no as appropriate.</p>

Supporting Documents:

<input type="checkbox"/> 1. Upload Paper Protocol.	If applicable, upload your most recent Paper Protocol that corresponds with this Muse protocol.
<input type="checkbox"/> 2. Upload Approval Letter.	If applicable, upload your most recent Approval Letter that corresponds with this Muse protocol.
<input type="checkbox"/> 3. Upload any Supporting Documents.	Upload any necessary supporting documentation or evidence: <ul style="list-style-type: none"> • personnel laboratory specific training or certifications • BSC or CFH certificates • PRCU SOP • Laboratory SOP • Procedural examples or photographic examples. • Grant Requirements, etc.

Your Protocol's Homepage:

1. Add PI Proxy:
 - a. If you have a Co-Investigator or are entering the protocol on behalf of the PI, assign that person/yourself as a PI Proxy so that they may submit any protocol modifications and receive protocol notifications along with the PI.

2. Manage Related IRB Studies and IACUC Protocols:
 - a. Click **Manage Related IRB Studies** to input information about your Human Clinical Trial.
 - b. Click **Manage IACUC Protocols** to input information about your Animal Use Protocol.
 - i. Note: this will allow the Animal Use protocols and Safety protocols to link in Muse.

Reminders:

- Save often and use the % to search, especially for room numbers.
- **If you cannot find an agent** (biohazard, chemical, etc.), email the Biosafety Officers (BSOs) to have the agent added in Muse (kaloyd@uams.edu and dmilke@uams.edu).

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- Also email the BSOs if you have any questions or need contact information for BSC Certifications.