Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
	Antidyskinetic Therapy for PD
Principal Investigator:	
Date of Action:	11/18/2021

The animal protocol review was considered by the Committee and the following decisions were made:

The protocol was approved.

<u>NOTE:</u> If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see <u>https://researchintegritv.asu.edu/animals/training.or contact</u> Research Support Services within DACT at

Additional requirements:

Cc:

IACUC Office IACUC Chair

\Box This protocol requires that Research Support Services group wit	hin DACT provide supervision for the
first time a procedure is conducted. Contact	chedule.
\boxtimes This protocol indicates that there are surgical procedures. A sur	reical checklist may be required to be
submitted to Research Support Services within DACT	prior to starting surgeries.
$oxed{intermat}$ Other requirements: IBC approval is required before work with	n biohazardous materials may begin

Total # of Animals: Species:	3 NHP	Pain Category: D	
Protocol Approval Period:	11/18/2021 - 11/17/2	024	
Sponsor: ASU Proposal/Award #: Title:			
Signature: IACUC Chair or	Designee		Date: 11/30/2021

Obtained by Rise for Animals. Uploaded to Animal Research Laboratory Overview (ARLO) on 08/15/2023

IACUC Use Only	IACUC Protocol #: 22-1886R
Date: 10/14/21	BC RSC Chem

ANIMAL USE PROTOCOL ARIZONA STATE UNIVERSITY INSTITUTIONAL ANIMAL CARE AND USE COMMITTEE (Revised February 2021)

Read "Instructions for Submitting the ASU Animal Use Protocol" before completing. Upon approval, this protocol will become a public record so follow instructions carefully.

PROJECT/PROGRAM TITLE: Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent

Antidyskinetic Therapy for PD

SPECIES REQUESTED: Cynomolgus macaque (Macaca fascicularis)

I. PERSONNEL INFORMATION

A. A single member of the university faculty and/or Principal Investigator (PI) is considered the responsible individual.

	NAME:		TITLE:	Director
	AFFILIATION:	ASU-Banner Neurodegenerative Disease Research Center	Office Phone #	
	Cell Phone #:		E-Mail:	
B.	Additional contact	, if any, for IACUC business		
	NAME:		TITLE:	Primate Lab Supervisor
	AFFILIATION:		Office Phone #	
	Cell Phone #:		E-Mail:	
C.	Protocol Type			
	Non-funded re	esearch		
	Internal Fundi	ng		
	Account Num	ber:		
	External Fund	ling (Grant/Contract)		
	Granting Ager	ncy:	Deadli	ne:
	Co-Investigate	or(s):		
	Proposal Title			
	ASU Proposa	or Award #		

If, ASU proposal or award number is not provided, attach a copy of the complete proposal or grant document.

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- Teaching Course Number and Title:
- D. Protocol Status:

🛛 New

- Renewal—Previous Protocol #:
- Revision—Previous Protocol #:
- E. Do you plan to use Department of Animal Care & Technologies (DACT) personnel and resources? If yes, describe the support needed? (If this use is new or an expansion of previous use, contact the DACT well in advance of need). Yes, surgical anesthesia and post-op monitoring, as well as standard husbandry and clinical care.

II. PROJECT DESCRIPTION AND PROGRAM REQUIREMENTS

The Institutional Animal Care and Use Committee (IACUC) is composed of both active animal users and lay persons. Regardless of background, each member has a vote, so it is particularly important that the language of the application be understood by all. This applies to all sections of the application, but it is especially important that the goals and justifications of the proposed research be spelled out in the clearest possible terms. NOTE: Upon approval, this protocol will become a public record, so do not disclose proprietary information.

A. Provide a brief (300 words or less) synopsis in <u>NON-SCIENTIFIC TERMS</u> of proposed research.

Symptomatic treatment of individuals with Parkinson's disease (PD) includes dopamine (DA) replacement therapies, with the drug levodopa, the gold standard and most potent pharmacological therapy to date. Long-term levodopa therapy is associated with motor complication side-effects including levodopa-induced dyskinesias (LID) which causes uncontrolled, involuntary movements. Selectively targeting calcium channels can provide potent long term prevention of LID even with extreme doses of daily levodopa. We plan to investigate the potential of a viral vector that targets calcium channels in non-human primates, the 'gold standard' PD model that allows for reversal of mild-to-moderate LID, and prevent escalation of LID severity with increasing doses of levodopa.

B. PLANNED USE OF ANIMALS. Begin with a clear statement of purpose and briefly provide background information and references to previous work (especially if this is a renewal protocol). Include a clear description of the experimental design for all animal experiments planned and explain why the experiments must be performed. It is critical that for each procedure you provide a detailed sequence of events that effectively describes what happens to the animals from acquisition to euthanasia (if applicable). As the focus of the IACUC protocol is on animal use, do not simply cut and paste research objective statements from grant proposals. Flow charts, diagrams or tables are strongly recommended for complicated experimental designs. State how the research is expected to benefit the human community, the animal community, and/or society as a whole. Details regarding surgical procedures, drug treatments, and field techniques are not necessary, as they will be addressed later in the form.

Background: Eliminating levodopa-induced dyskinesias (LID) is a significant unmet need in Parkinson's disease (PD) therapy. There are currently no FDA approved drug treatments for LID, yet up to 90% of individuals with PD develop this side-effect. The L-type calcium channel CaV1.3 is a target of interest for LID prevention. Towards this end, the pathognomonic loss of striatal dopamine (DA) in PD results in dysregulation and overactivity of striatal CaV1.3 channels leading to synaptic pathology, including the loss of dendritic spines on striatal spiny projection neurons that appears to be involved in LID. While initial studies performed by our colleagues at delivering CaV1.3 channel antagonists reduced LID dose-dependently, the

effects were partial and transient. To provide unequivocal target validation, free of pharmacological limitations, then developed a rAAV-CaV1.3-shRNA to provide continuous, high potency, target-selective, mRNA-level silencing of striatal CaV1.3 channels. In rats, using software and RNAscope in situ hybridization (ISH) with custom RNA target probes obtained from the striate software and RNAscope in situ hybridization (ISH) mRNA silencing, ISH revealed an average of 85% less mRNA transcript in the region of the striatum stained positive for the reporter protein GFP in rAAV-CaV1.3-shRNA rats compared to rAAV-Scr-shRNA rats. The range

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of mRNA silencing was 43.6% to 99.1% indicating that even partial silencing (or normalizing) of CaV1.3 channel over activity in the parkinsonian striatum is capable of completely preventing LID induction.

then examined whether genetic silencing of these dysregulated calcium channels could prevent LID induction in previously levodopa naïve parkinsonian rats and/or whether it could reverse these abnormal behaviors in parkinsonian rats already expressing a severe LID phenotype. In their 'LID prevention studies' they found that gene level silencing of striatal CaV1.3 channels in severely parkinsonian rats, prior to the introduction of levodopa: 1) can provide uniform and complete protection against the induction of LID, and 2) that the antidyskinetic benefit is sustained over time and even with high doses of daily levodopa. In our 'LID reversal studies' we observed that AAV-mediated CaV1.3 silencing in parkinsonian rats with already established LID could ameliorate these behaviors, however: 1) a one-week drug withdrawal 'drug holiday' appeared beneficial and/or necessary, and 2) the antidyskinetic effect was variable in rats expressing protracted, high levels of LID that accompanies chronic high dose levodopa. Importantly this approach did NOT interfere with motor benefit of levodopa. Gene delivery resulting in striatal CaV1.3 silencing provides some of the most profound antidyskinetic benefit reported to date. If these findings can be translated into a clinical application with a similar magnitude, this would provide a much-needed breakthrough in treatment of individuals with PD and would allow the most powerful antiparkinsonian therapy ever identified to work unabated through the duration of the disease. Since the non-human primate model is the gold standard for preclinical testing and this shRNA has not been used in NHP before, we will extend these proof-of-principle studies by examining the antidyskinetic utility of rAAV-CaV1.3shRNA in parkinsonian cynomolgus macagues. In this experiment, we propose to examine the capacity of striatal CaV1.3 channel silencing to prevent induction of LID in this model.

Experimental Design: 3 cynomolgus macaques (M, ≥20 years old) will be transferred from the

where these animals have previously been rendered parkinsonian via intracarotid and systemic MPTP administration. Their level of impairment has been stable for at least one month after the last MPTP injection and is not expected to progress. All three animals exhibit mild to moderate motor impairments but are able to eat and drink and move about their cages on their own. Once transferred to ASU, all subjects will be retrained on the hand reach task, and be rated on the clinical rating scale. The animals will have baseline serum, plasma, & cerebrospinal fluid (CSF) collected (during the same session of sedation when possible). Animals will then receive pre-op MRI for surgical targeting. We will then deliver bilateral injections of rAAV-Cav1.3-shRNA (n=1) or rAAV-Scr-shRNA (n=2), stereotaxically into the putamen. Beginning seven weeks post-injection, animals will begin behavioral testing (hand reach task and CRS) with and without levodopa/carbidopa treatment every four weeks. Eight weeks after vector injection all monkeys will begin twice daily (M-Fr) treatment with levodopa/carbidopa (20 mg/kg levodopa). 20 weeks after injection, the levodopa dose will be increased to 30 mg/kg, and 26 weeks after injection the dose will be increased to 40 mg/kg. Serum, plasma, & CSF will be collected 4, 8, & 16 weeks post injection and at sacrifice. Beginning eight weeks post-injection and continuing every two weeks thereafter, animals will be videoed in their home cage or in a separate designated cage at multiple time points after their morning levodopa/carbidopa administration for LID scoring. Animals will be sacrificed 30 weeks post-injection and their brains collected. All collected tissue will be sent to

Procedures:

MRI Scanning (one or two times): Stereotaxic intracranial injections are performed under intraoperative MRI guidance. Preop MRIs will be <u>performed</u> on MRI scanner at the

After transportation to animals will be anesthetized with ketamine (3-10 mg/kg, IM) and either dexmedetomidine (0.03 mg/kg, IM) or midazolam (0.05-0.5 mg/kg) and maintained with gas anesthesia (e.g., isoflurane, sevoflurane) or booster injections of ketamine (1.5 mg/kg, IM) and dexmedetomidine (0.015 mg/kg, IM). Animals will be intubated to maintain a stable airway. Cetacaine spray (200 mg, topical) may be applied to the throat to assist with intubation. Animals will be placed in an MRI compatible stereotaxic frame and MRI-opaque fiducial markers will be placed around the skull for neuronavigation registration. T1 and/or T2-weighted images will be obtained. The locations of the fiducial markers will be permanently marked with a tattoo dot on the skin using a commercial tattoo marker. The MRI scan time is approximately 20 minutes, sedation is expected to last ≤1 hour. Following the procedure,

dexmedetomidine/midazolam may be reversed with atipamezole (0.15-0.3 mg/kg, IM) or flumazenil (0.025 mg/kg, IV), respectively. As MRI registration is essential for accurate surgical guidance, if in the opinion of the surgeon the MR images prove inadequate (due to animal movement, fiducial placement, or other confounding factors), up to one repeat MRI may be performed.

Fine Motor Skills Test (Hand Reach Task) (six times per week, every four weeks, beginning seven weeks post-vector injection): Animals will be tested for fine motor performance in both upper limbs using a hand reach task (HRT). Animals will be transported to a modified testing cage from their home cage and presented with a 3x2 well matrix plexiglass testing board. Four pieces of food will be placed within the wells for each trial, and time will be recorded for how long it takes the animal to retrieve them. The board is configured so that only one limb will be tested at a time. Animals will undergo 10 trials per limb, with each trial being alternated between the left and right limbs. Each animal will be tested by the same investigator. Investigators will be blinded to the animal's treatment groups. If necessary, food restriction will be used to encourage animal participation in this activity.

Clinical Ratings Scale (at least once per month): Animals will be evaluated with the Clinical Ratings Scale (CRS), a collection of clinical observations, scored and graded in order to determine the severity of an animal's overall impairments (see below). The ratings scale contains scoring in the categories of posture (0-3), gait (0-5), bradykinesia (0-5), balance (0-3), tremor in left and right arms (0-3 for each), gross motor skills in left and right arms (0-4 for each), defense reaction (0-2) and freezing (0-2). The combined scores will determine the impairment level of the animal, a score of 0 indicates a normal animal and a score of 12 or higher indicates parkinsonism. Individual scores in each category can be assessed separately as well in order to determine the overall health and well-being of an animal. Occurrence of dyskinesia, psychological disturbances and vomiting will also be noted. In the past, animals that have been rendered parkinsonian with MPTP have remained stable and been able to eat and drink on their own. However, if any animals begin to display new neurological deficits or other clinical signs that may impact their health and well-being, including a posture score of 3, gait score of 3 or more, bradykinesia score of 3 or more, or gross motor skills score of 3 or more in both arms, they will be referred to the veterinary staff for evaluation and treatment, if necessary.

Posture (0-3)	Gait (0-5)	Bradykinesia (0-5)	Balance (0-3)	Tremor (0-3)	Gross Motor Skills (0-4)	Defense Reaction (0-2)	Freezing (0-2)
0-Normal, upright posture 1-Mildly stooped, neck and shoulders slightly curved 2-Notably stooped 3-Face down, unable to sit	0-Normal, smooth movements 1-Mildly impaired 2-Moderately impaired, some stumbling, abnormal footing (crossover) 3-Severely impaired, stumbling, bradykinetic 4-Severely impaired, loss of balance, freezing 5-Incapable of movement	0-Normal, prompt, brisk, plentiful movements 1-Mild slowness, lesser overall movements than normal 2-Moderate slowness, increasing poverty of movement 3-Moderate slowness with freezing, few and labored movements 4-Severe slowness with freezing, few and labored movements 5-Unable to ambulate	0-Normal 1-Diffculty standing, holding onto cage bars; holds cage bars intermittently but can stand alone 2-Unable to stand without holding onto cage bars 3-Unable to sit without leaning; completely unable to stand	0-Absent 1-Small amplitude and/or infr 2-Large amplitude and/or frequent 3-Occurs constantly; interferes with normal behavior	0-Normal 1-Mild impairment, slight clumsiness 2-Moderately impaired, difficulty manipulating small objects, often drops food reward 3-Does not readily use arm to reach for food; can use to ambulate 4-Cannot use arm/hand for feeding or ambulating	0-Normal, aggressive, comes to front of cage, shakes bars, threatens 1-Strong facial threat but does not move 2-Minimal or no response	0-No freezing 1-Notable freezing; freezing readily broken with outside stimulation 2-Large periods of time frozen; interferes with ambulation

LID Video Recording (six times per day, every two weeks, beginning eight weeks post-vector injection): Animals will be video recorded in their home cage or transferred to a designated video cage on the day of recording. They will remain in the designated video cage until the final recording session of the day. Animals will be recorded for 3 minutes prior to levodopa/carbidopa dosing, and then again for 3 minutes at 20, 70, 120, 170, and 220 minutes post levodopa/carbidopa dosing.

Levodopa/Carbidopa Dosing (twice per day, M-Fr, beginning eight weeks post-vector injection): Levodopa/Carbidopa in a combined pill (4:1 levodopa/carbidopa ratio, 20-40 mg/kg levodopa/5-10 mg/kg carbidopa) will be crushed and mixed with a food treat (fruit, jelly, honey, etc.) and given to the animal in a paper cup in the morning (9-10 am) and afternoon (4-5 pm).

Euthanasia (once): The animal will be anesthetized with ketamine (10 mg/kg, IM), xylazine (2 mg/kg, IM) or midazolam (0.05-0.5 mg/kg), and either hydromorphone (0.2 mg/kg, IM) or morphine (1 mg/kg, IM), followed by gas anesthesia (e.g., isoflurane, sevoflurane) or additional drug delivered IV [ketamine (\leq 20 mg/kg, IV), xylazine (\leq 4 mg/kg, IV), and either hydromorphone (\leq 0.4 mg/kg, IV) or morphine (\leq 2 mg/kg, IV)], if needed to achieve a surgical plane of anesthesia. Once a surgical plane of anesthesia is achieved as verified by lack of response to toe/finger pinch, palpebral reflex, and corneal reflex, the thoracic cavity will be opened, heparin (5,000 IU, IC) will be injected into the left ventricle of the heart and the animal will be euthanized via transcardial perfusion of 0.9% saline (1-2 L) followed by 4% buffered formaldehyde (1-2 L). If perfusion cannot be performed for any reason,

animals will be anesthetized with ketamine (10 mg/kg, IM) followed by a pentobarbital-containing euthanasia solution (86-120 mg/kg, IV).

C. RATIONALE FOR INVOLVING ANIMALS AND THE APPROPRIATENESS OF THE SPECIES AND NUMBER USED. Keeping in mind the principles of the "3 R's" (Refinement, Reduction, and Replacement), answer the following:

1. Why must live vertebrates be used in this study?

Our laboratory's experiments are aimed to help human patients with neurodegenerative diseases. Preclinical research requires a model that can inform about the applicability of diagnostic tools as well as the potential complications of its utilization. To the best of our knowledge, in vitro and computer models are still not able to give us enough information for clinical projection in complex neurodegenerative diseases such as PD.

2. Why are you using the requested species rather than other species?

Cynomolgus macaques were chosen because their brains closely resemble that of a human, in comparison to a smaller or less complex species. MPTP lesioned primates provide a useful model for mimicking human PD and are considered the best pre-clinical species for testing cell-based therapeutics. Rodents do not effectively mimic the complex degeneration in nigrostriatal circuitry necessary to appropriately test PD therapies pre-clinically.

3. What is the rationale supporting the numbers of animals proposed? Typically, a power analysis should be performed to support the proposed sample sizes. A table depicting the number of animals to be used is required.

This protocol will include an N of 3 animals, which includes 1 animal in group 1 and 2 animals in group 2. The results from these 3 animals will be combined with the results of the 9 animals previously completed a to obtain our total of N=6 per group for statistical analysis.

All LID and CRS behavioral data will be analyzed with non-parametric statistics including Mann-Whitney Utest (for between-subject contrasts) and Friedman or Kruskal-Wallis with Dunn's multiple comparison test (for with-in subjects tests). Post-mortem data (e.g.: TH+ cell number) and behavioral tests that provide parametric data (e.g.: HRT) will be analyzed by t-tests or multifactorial ANOVAs with appropriate post-hoc analyses after testing for normal distribution. Total animal number and group sizes have been determined by power analysis based on data from our previous experience with these models and using the software G*Power 3.1.9.2 program. The number of animals proposed (n=6 per group) allows us to be well-powered to detect a 15-20% change, ($\beta > 0.8$, $\alpha \le 0.05$).

Group	Treatment	Total N=	N= as of 10/2021	N= in this protocol
1	rAAV-Cav1.3-shRNA	6	5	1
2	rAAV-Scr-shRNA	6	4	2

4. What refinements, if any, have been made to reduce the number of animals used and the potential detrimental effects on the study animals?

We feel the number of animals requested is the minimum necessary to achieve the aims of this study. Furthermore, we have taken every precaution to avoid pain and discomfort in our animals. The intracranial injections proposed will be conducted under general anesthesia with proper perioperative pain management and postoperative monitoring and care.

III. EMERGENCY CONTACT

A. Who should be contacted in case of an animal emergency? Note: This information will be redacted if this protocol is requested as a public document.

Name: Office Phone # Home Phone # Cell Phone #:



IV. DUPLICATION AND ALTERNATIVES PLEASE READ ALL INSTRUCTIONS.

The Animal Welfare Act requires that you document your justifications with data from two or more sources. <u>One</u> source must be a set of searches of a relevant database: name the database searched, the keyword and keyword combinations searched, the date the search was performed and the date range searched. The second source can be a set of searches of a second relevant database, or consultation with a laboratory animal science veterinarian, or courses/meetings/consultations with qualified personnel. Sufficient documentation, such as the consultant's name and qualifications and the date and content of the consult, should be provided to the IACUC to demonstrate the expert's knowledge of the availability of alternatives in the specific field of study. Examples of appropriate databases to search include PUBMED, Web of Science, or Animal Welfare Information Center (AWIC – recommended for USDA-covered species https://www.nal.usda.gov/awic/databases).

A. Provide the following details for the most recent literature search used to explore for <u>duplicative research</u>. (The literature search documents that the research will not unnecessarily duplicate previous research). **Teaching protocols do not need to conduct this search**.

Date that search was conducted (*Must be within 60 days of the IACUC review date*): 10/14/2021 Database(s) used: ALTBIB, PUBMED Publication years covered by the search: 2000 - present Keyword combinations used: Nonhuman primate, levodopa-induced dyskinesia, CaV1.3

B. Provide the following details for the most recent literature search used to explore for <u>alternatives to animal use</u> and <u>alternatives to painful procedures</u>. Alternatives should be considered for any aspect of the protocol that may cause more than momentary or slight pain or distress to the animal. Alternatives to be considered include those that would: 1) refine the procedure to minimize discomfort that the animal(s) may experience; 2) reduce the number of animals used overall; or 3) replace animals with non-animal alternatives (e.g., computer models or tissue culture). All protocols (research and teaching) MUST conduct this search.

Date that search was conducted (*Must be within 60 days of the IACUC review date*): 10/14/2021 Database(s) used: ALTBIB, PUBMED Publication years covered by the search: 2000 - present Keyword combinations used: Parkinson's disease, animal model, nonhuman primate Nonhuman primate, intracranial injection alternative

C. Results of literature search for alternatives: Comment on the application(s) of any identified alternatives (found with your search terms, including how these alternatives may be or may not be incorporated to modify a procedure to either lessen or eliminate potential pain and distress. All protocols must complete this section and must describe how the literature search results relate to painful procedures and alternatives to animal use. You must include sufficient information for the IACUC to determine that a reasonable, good faith effort was made to determine the availability of alternatives. If the search identified any alternative methods (ones that could be used to accomplish the goals of the animal use proposal), you must clearly explain and justify why this alternative cannot be used.

For instance, if your search terms retrieved eight publications, summarize how many of those described alternatives to painful procedures and the use of animals. No studies have been published examining the role of CaV1.3 in preventing levodopa-induced dyskinesias in an NHP model of PD. While this vector has been studied in a rat model of PD, the NHP models of PD better mimic what is seen in the human brain, increasing the applicability of the results to human patients. Additionally, the brains of rodents are less complex than that of NHPs and humans. No alternatives to intracranial injection were found for delivering vector to the putamen.

D. Describe any other procedures (e.g., participation in meetings, review of journals) that are used to explore and evaluate alternatives: The PI, lab manager, post-docs, and graduate students regularly attend national meetings

and discuss recent updates in technology and methodology for these experiments with colleagues. Additionally, they remain up to date with the scientific literature on new and alternative procedures.

- E. Does this research replicate previous work? (Your answer will be based in part on the literature search above.)
 - \boxtimes No. Proceed to section **VI**.
 - ☐ Yes. Explain why the replication is necessary:
 - Not applicable. This is a teaching protocol.

V. CATEGORY OF PAIN OR DISTRESS

For non-USDA covered species, answer question A only. For USDA covered species, answer question B only. USDA covered species are all mammals EXCEPT laboratory mice and rats bred for research. All other rodents, including wild mice and rats, are covered.

A. Do the procedures in this protocol have the potential to involve more than slight or momentary pain or distress that will **NOT** be relieved with anesthetics, analgesics, tranquilizer drugs, or other method for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia)?

If yes, describe and justify:

B. Using the table below, list all USDA covered species to be used in the proposed study and indicate the number of animals to be used under each USDA pain category. For an animal undergoing multiple procedures, include the animal under the highest level of pain/distress expected for that animal.

	Number per UŞDA Category*			Total number of	
USDA Covered Species	В	С	D	E	animals requested
Cynomolgus macaque			3		3

*USDA PAIN CATEGORIES: (see <u>http://researchintegrity.asu.edu/animals/forms</u> for a more complete description of the below categories)

<u>Classification B:</u> Includes animals that are used solely for breeding or are being acclimatized or held for use in teaching, testing, experiments, research, or surgery but have not yet been used for such purposes.

<u>Classification C:</u> Includes the use of animals in procedures involving no, momentary, or slight pain or distress (e.g., non-invasive parenteral drug delivery, peripheral blood collection, euthanasia, short-term manual or chemical restraint, toe clipping).

<u>Classification D:</u> Includes the use of animals used in procedures that could cause pain or distress but appropriate anesthetics, analgesics, and/or tranquilizing drugs or other methods for relieving pain or distress are used (e.g., surgery, perfusion, administration of irritating chemicals, humane endpoint euthanasia).

<u>Classification E:</u> Includes the use of animals in procedures that have the potential to involve pain or distress that will **not** be relieved with anesthetics, analgesics, tranquilizer drugs, or other method for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia).

VI. ASSURANCE:

The information contained herein is accurate to the best of my knowledge. I have carefully compared the proposed work with the current state of knowledge in this field by reviewing the literature and it is my professional opinion that the proposed work meets high standards of scientific merit. If the study involves pain and distress to the animal, whether or not it is relieved by anesthetics or analgesics, I have (1) reviewed the literature related to this work and have found no significant studies which could make this protocol <u>unnecessarily</u> duplicative, and (2) considered alternatives to animal use and found none available, as described above. Procedures involving animals will be carried out humanely and all procedures will be performed by or under the direction of trained or experienced persons. Any revisions to animal care and use in this project will be promptly forwarded to the Institutional Animal Care and Use Committee for review. Revised protocols will not be used until Committee clearance is received. The use of alternatives to animal models has been considered and found to be unacceptable at this time.

The principal investigator, by signing below, and the IACUC recognize that other medications may be given to the animals for veterinary care purposes. This includes the humane euthanasia of animals in uncontrollable pain or distress as determined by the Attending Veterinarian or the Clinical Veterinarian acting for the Attending Veterinarian. However, the veterinarians will make all efforts to contact and discuss the case with the Principal Investigator or designee prior to making a unilateral decision.

	10/14/21
Principal Investigator –Print	Date
	10/1 1/01
	10/14/21
Principal Investigator Signature	Date

NOTE: Principal investigators must submit a current curriculum vitae or biosketch that reflects their most recent pertinent experience.

PERSONNEL CHART

ASU requires that all personnel engaged in animal research or teaching be qualified through training or experience in order to conduct the work humanely. The IACUC requires the following training:

- Level I Basic Required of ALL participants (must be renewed every 4 years)
- Level II Species-Specific Required for each participant that will have direct contact with that species (must be renewed every 4 years)
- Level III Hands-on Training Required to perform specific procedures independently. Training can be received from DACT staff or PIs can have a member of their lab approved by DACT to be a Certified Trainer who can then train others. Simply being Level III certified does not allow the training of others. A Level III Certification form must be submitted to the IACUC office by the person providing the training within 5 days of the training.

You can access the training modules at <u>https://asu.co1.gualtrics.com/jfe/form/SV_b2b2XRXRRs1309f.</u> See the IACUC web site (<u>https://researchintegrity.asu.edu/animals/training</u>) for more information on training and Level III forms.

* Procedures other than husbandry, handling, or behavioral testing MUST be performed under supervision unless the person is Level III certified to conduct the procedure independently. Personnel are not Level III certified until the IACUC has reviewed and approved the Level III training documentation. The PI is responsible for ensuring that personnel who are not Level III certified are supervised at all times.

Name Title direct supervision? submission? species) Conf Intracranial surgery, Intracranial surgery, Intracranial surgery, blood/CSF collection, 7/2021 Intracranial surgery, Intracranial surgery, Blood/CSF collection, MRI, administration of any medications, and 1 Intracranial surgery, PI Intracranial surgery, All I I	Name

For each individual, describe the individual's years of experience with all listed species and procedures they will be conducting under this protocol. For procedures for which they are not yet trained, but will likely be trained to do during the activity period of this protocol, provide a description of who will provide such training: Drawn has 37 years' experience conducting research with nonhuman primates and is experienced with all procedures in this protocol.

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DETAILED USE OF ANIMALS

This section must be completed for each species used.

(additional Detailed Use of Animals forms can be found at https://researchintegrity.asu.edu/animals/forms)

Common Name: Cynomolgus macaque

Scientific Name: Macaca fascicularis

I. ANIMAL INFORMATION

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- A. Is this a threatened or endangered species?
 - No. Proceed to section I. B.
 - Yes. Describe why this work must be done on this species and why the project will not have a significant negative impact on the species:
- B. Maximum # of animals to be used over the 3-year life of the protocol: 3
- C. Sex: M Age or Weight Range: ≥22 years
- D. Source (e.g., commercial, in-house breeding, captured from wild): Transfer from
- E. List all labs and/or rooms outside of the ASU centralized vivaria where you intend to keep or use live animals in connection with the animal use covered under this protocol. This list is for IACUC information to assure each location is inspected semi-annually. Listing rooms here does not assure approval of this space for use.

Building	Room #	Max Length of Stay	Method of Transport	Purpose
			NHP cage inside DACT truck	MR Imaging

F. If you use DEA-controlled substances, list the location where they are stored (building and room number). If you acquire controlled substances from DACT for same day use, state this. The IACUC is required to inspect all controlled substance storage locations semi-annually. Controlled substances will be stored in Drooffice

II. MAJOR CATEGORIES OF USE

- A. Will animals be immunized solely for the production and harvesting of antibodies to be used in vitro rather than as a vaccine study?
 - No. Proceed to section II. B.
 - Yes. Complete the following table.
 - Injection:

Volume of injectate	Adjuvant	Route	Min. Frequency	Max. # of injections

Collection: If terminal, check here D otherwise complete the following.

Route	Max. Volume	Min. Frequency	Max. # of collections	

- B. Will tissues, blood, or other body fluids be harvested (other than for antibody production)?
 - No. Proceed to section II. C.
 - Yes. Will tissues, blood, or other body fluids be collected post-mortem only?
 - Yes. Proceed to section II.C.
 - No. Complete Appendix 1: Antemortem Specimen Collection.
- C. Will animals be food restricted (calorically or specific constituents) other than for surgical procedures? No. Proceed to section II. D.

Yes. [note: restriction paradigms exceeding a single 24-hr period must follow the ASU IACUC Standard Institutional Guideline for Food and Water Restriction available at https://researchintegrity.asu.edu/index.php/animals/procedures-library-and-guidelines

1. What are the restriction parameters? Provide scientific justification and include the length of restriction. Some animals are initially resistant to performing the hand reach task (HRT). In order to provide increased incentive for these animals, we will offer different food treats (cereal, marshmallows, raisins, peanuts, etc.) to discover the animals' preferences (stage 1). If that is unsuccessful after one week, we will postpone daily feeding of the normal food allotment until after the animal has been tested (stage 2). If that is unsuccessful after one week, we would like to temporarily restrict the number of food biscuits fed to the animals (stage 3). Adult cynomolgus monkeys normally receive 6-12 biscuits per day and a half a fruit or vegetable, food restricted animals will receive a minimum of half their normal allotment of biscuits based on veterinary recommendation, along with a half a fruit or vegetable. Food restriction during the initial HRT training period will last a maximum of one month. Previous animals that have been placed on similar food restriction have not become combative in our experience, but food restricted animals will be monitored and will be temporarily separated from their partners during feeding if necessary. This incentive program will apply to all animals participating in the HRT initial training and baseline. If an animal becomes reluctant to participate in the HRT at any time following surgery, they may be food restricted up to a maximum of one week (7 days) out of every four weeks during which time HRT performance will be recorded. A log of the HRT incentive program will be maintained.

2. How will you monitor for negative effects of food restriction (include information on how you will account for animal growth)?

Prior to the initiation of food restriction, the veterinary staff will be consulted on the animal's ideal body weight and current body condition score (BCS) on a 9-point scale (see below). Animals with a BCS of 3.5/9 or less will not be placed on food restriction. During food restriction, the animal's body weight will be monitored every week or twice per week if body mass loss exceeds 8% of starting mass. An animal that loses 10% or more of its body weight from the start of food restriction or whose BCS drops to 3.5/9 or below will be removed from food restriction and will return to stage 2 (postponement of feeding full daily ration until after testing). Food restriction will last up to a maximum of one month during the initial HRT training period and up to a maximum of one week (7 days) out of every four weeks during the post-surgery assessment period. The same body weight and BCS restrictions will apply during this time.

- D. Will animals be water restricted?
 - No. Proceed to section II. E.
 - Yes. [note: restriction paradigms exceeding a single 24-hr period must follow the ASU IACUC Standard Institutional Guideline for Food and Water Restriction available at https://researchintegrity.asu.edu/index.php/animals/procedures-library-and-guidelines
 - 1. What are the restriction parameters? Provide scientific justification and include the length of restriction.
 - 2. How will you monitor for negative effects of water restriction (include information on how you will account for animal growth)?
- E. Will animals be exposed to trauma, injury, burning, freezing, electric shock, UV radiation, magnetic fields, lasers, loud noise, or other physical agents that might cause distress?
 - No. Proceed to section II. F.
 - \boxtimes Yes. List and justify each exposure.

Provide scientific justification: Magnetic Resonance Imaging (MRI): Due to the variability in NHP neuroanatomy, MR imaging is the best way to accurately target surgical injections within the brain. MRI scans involve strong magnetic fields, and precautions are made to ensure that no incompatible metals are present in the room during the scan. Noise levels inside an MRI machine typically vary from 65 to 95 dB, and intermittent spikes of ~110 dB may be produced. MRI scans are performed under anesthesia, and ear protection using ear plugs or gauze/cotton will be placed in the animal's ears to prevent damage and mitigate distress.

- F. Will animals be exposed to environmental stress (e.g., non-natural temperature exposure, prolonged physical restraint, forced exercise)?
 - \boxtimes No. Proceed to section II. G.
 - Yes. List and scientifically justify each exposure.
- G. Will animals undergo surgery?
 - No. Proceed to section II. H.
 - Yes. Complete Appendix 2: Surgical Procedures.
- H. Will any animals have a device (e.g., thermocouple, cannula, electrode) that extends chronically through the skin? No. Proceed to section II. I.
 - Yes. Describe wound management measures to minimize chances of infection around the device where it penetrates the skin:
- I. Will individuals of a social species (e.g., most rodents) need to be housed singly at any time?
 - No. Proceed to section II. J.
 - Yes.
 - 1. What would be the maximum duration that an individual would be singly housed? Provide scientific justification for singly housing for this duration: Animals will be pair housed when possible. However, because of the relatively small number of animals involved, suitable pairing partners may not be available. If necessary, single housing will be determined in conjunction with the veterinary staff and will continue until a suitable pairing partner becomes available or the experiment concludes. Animals that will undergo HRT will be permanently exempt from the establishment of new social housing pairs (contact or mesh) beginning with baseline behavioral data collection. An injury to the arm, hand, or fingers can severely affect the HRT results and the establishment of new social housing pairs is a common cause of injuries. Animals in established pairs will be allowed to remain partnered. However, if injurious fights occur between established pairs, the animals may be permanently separated based on consultation with the veterinary staff.
 - 2. Singly housed animals should receive additional enrichment. Describe what enrichment will be provided or scientifically justify why additional enrichment cannot be provided: Animals will be housed in a room with other conspecifics and have access to visual, olfactory, and vocal/auditory contact. All animals are also provided a variety of enrichment items including manipulanda and destructibles; these may be increased in number or variety for singly housed animals as determined by the veterinary staff on a case-by-case basis.
- J. Will animals need any special husbandry considerations, including but not limited to altering standard cage type, cage change frequencies, housing temperature, or lack of enrichment?
 - No. Proceed to section II. J.
 - Yes. Describe special procedures and provide scientific justification:
- K. Will animals be transported off campus (e.g., to/from the field, or between institutions) in a vehicle other than one owned by the DACT?
 - No. Proceed to section II. L.
 - Yes. Describe details (e.g., vehicle to be used, destinations, and driven by whom), read the IACUC SIG Off-campus Transport of Animals by Laboratory Personnel, and complete and submit with this protocol the Assurance to Abide by the Requirements for Transporting Live Animals:
- L. Will any work be conducted in the field (this includes field experiments or the capture of animals to be used in laboratory experiments)?
 - \boxtimes No. Proceed to section II. K.
 - Yes. Complete Appendix 3: Field Research.
- M. Will any animals need to be individually identified?

- No. Proceed to section III.
- Yes. Describe the marking technique to be used, why that technique was chosen, how it will be performed, and on what age range of animals?

Animals will be tattooed with an identification number on their chest or inner thigh. Animals either have the tattoo upon arrival or are tattooed while under sedation by DACT staff during quarantine. Touch ups may be done while sedated/anesthetized (e.g., for TB testing), and the hair in the region is shaved as needed to maintain visibility of the tattoo. This identification method is the most widely used means of permanently marking macaques.

III. CHEMICALS AND OTHER POTENTIAL HAZARDS

(If you answer yes to any of the following questions, this information may be forwarded to another oversight unit to aid you in assuring safe practices. Approval by these units or additional training may be required prior to using any of these materials)

- A. Will drugs or chemicals be used with animals?
 - No. Proceed to section III. B.
 - Yes. For each drug or chemical, list the agent, dose, route, purpose, and grade in the table below:

<u>Agent</u>	<u>Do e</u>	<u>Route</u>	Purpo_e	Frequency	Pharmaceutical grade (Y/N)?	Is this a DEA controlled substanc e (Y/N)?
Atipamezole	0.15-0.3 mg/kg	IM	Dexmedetomidine reversal	As needed	Y	Ν
Atropine	0.02-0.05 mg/kg	IM	Reduce respiratory secretions and prevent bradycardia	As needed	Y	N
Betadine	N/A	Topical	Topical disinfectant	As needed	Y	N
Bupivacaine	1-2 mg/kg	SC	Analgesia	Once during closure	Y	N
Buprenorphine Sustained release	0.2 mg/kg	SC	Analgesia	Once post- op	Y	Y
Cefazolin	20-25 mg/kg	IV or IM	Antibiotic	Every 2-4 hours intra- op, as needed	Y	N
Cephalexin	20-30 mg/kg	PO	Antibiotic	Twice daily, as needed	Y	N
Cetacaine Spray (Benzocaine 14%, Butamben 2%, Tetracaine 2%)	200 mg	Topical	Anesthesia	As needed for intubation	Y	N
Chlorhexidine	N/A	Topical	Topical disinfectant	As needed	Y	N
Dexmedetomidine	0.015-0.05 mg/kg	IM	Anesthesia	As needed	Y	N
4% Formaldehyde	1-2 L	IC	Perfusion	Once	Ν	N
Flumazenil	0.025 mg/kg	IV	Benzodiazepine reversal	As needed	Y	N
Gelfoam	Cut to size	Topical	Hemostasis/Seal surgical holes	As needed	Y	N
Heparin	5,000 IU	IC	Anticoagulant for perfusion	Once	Y	N

Hydromorphone	0 05 0 4 mg/kg	SC, IM, IV	Analge ia	A needed	Y	Y
Isoflurane	0.5-5%	Inhalation	Anesthesia	As needed	Υ	N
Isopropyl alcohol	70%	Topical	Topical disinfectant	As needed	Y	N
Ketamine	1.5-20 mg/kg	IM, IV	Anesthesia	As needed	Y	Y
Levodopa/Carbido pa (4:1 ratio)	20-40 mg/kg / 5-10 mg/kg	PO	PD therapeutic	Twice daily M-Fr	Y	N
Meloxicam	0.1-0.2 mg/kg	PO, SC	Analgesia	Once daily, a needed	Y	N
Melo icam Sustained release (10 mg/mL)	0 6 mg/kg	SC	Analge ia	Once po t op	Y	N
Midazolam	0.05-0.5 mg/kg	IM, IV	Sedative, anticonvulsant	As needed	Y	Y
Morphine	1-2 mg/kg	IM, IV	Analgesia	As needed	Y	Y
Ophthalmic ointment	Dab	Topical	Prevent corneal desiccation	As needed	Y	N
Pentobarbital- containing euthanasia solution	86-120 mg/kg	IV	Euthanasia	Once	Y	Y
Propofol	2-5 mg/kg Bolus 0.2-0.6 mg/kg/min CRI	IV	Anesthesia	As needed Continuous, as needed	Y	N
Saline or Lactated Ringer's Solution	5-15 mL/kg/hr	IV	Fluid replacement	Constant- rate infusion	Y	N
Saline	1-2 L	IC	Perfusion	Once	Y	N
Sevoflurane	1-8%	Inhalation	Anesthesia	As needed	Y	N
Sufentanil	0.25-2 µg/kg/hr	IV	Analgesia	Constant- rate infusion	Y	Y
Xylazine	2-4 mg/kg	IM, IV	Anesthesia	As needed	Υ	N

- 1. For each drug or chemical that is not pharmaceutical grade, indicate whether no pharmaceutical grade equivalent exists or provide scientific justification for using the non-pharmaceutical grade product. Formaldehyde is not available in a pharmaceutical grade, and is only used once in a terminal procedure.
- B. Does this project involve transgenic, knockout, or knock-in animals?

 \boxtimes No. Proceed to section III. C.

Yes. List the strains, any special care needs, and any expected clinical signs that are associated with the strain. Transgenic animals need to be covered by an IBC disclosure.

C. Does this project involve the use of biohazardous agents in animals (microorganisms, microbial toxins, recombinant DNA)?

No. Proceed to section III. D.

 $\overline{\boxtimes}$ Yes. List the agent, as well as concentration, dose, and route if applicable.

Agent	Concentration	Dose Route	ADMIN. USE ONLY
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or Animals.

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				ABSL	IBC # if Reg'd
rAAV Cav1 3 hRNA	1E10 ¹³ vg/mL	80µL	Intracranial bilateral injection into putamen	Pending	SPROTO2021 70
rAAV-Scr-shRNA	1E10 ¹³ vg/mL	80µL	Intracranial bilateral injection into putamen	Pending	SPROTO2021-70

- D. Does this project involve irradiation or the use of radiological material in animals?
 - No. Proceed to section III. E.
 - Yes. List the agent, dose, route, and purpose in the table below:

Agent	Dose	<u>Route</u>	Purpose
	-		

- 1. Provide the date of Radiation Safety Committee approval:
- E. Describe any health hazards to researchers and include a description on how the risk is mitigated or managed: Risk of bites, scratches, or Herpes B (Herpes B virus is not being used in animals but can be transmitted to personnel if there is an NHP bite/exposure). Risks are mitigated with the use of additional PPE as required by University policies (such as, but not limited to, Tyvek sleeves and double gloves), NHP primate certification, annual B Virus training (including Bite/Scratch policy), proof of 2 MMR vaccines or a measles titer, annual TB screening, and ear protection during MRI scans.
- F. Describe any health hazards to animals and include a description on how the risk is mitigated or managed: Zoonosis such as TB, measles, and flu are agents of concern that may spread from humans to monkeys. Before working with an NHP, all researchers are required to show proof of 2 MMR vaccines or a measles titer and annual TB screening. All people interacting with the monkeys are also required to wear a surgical mask to prevent the spread of these infections.

IV. DETRIMENTAL SEQUELAE

- A. Will animals possibly experience clinical signs intentionally or as a possible side effect of the study? No. Proceed to section V.
 - $\overline{\triangleleft}$ Yes. Complete the following.

res. Complete the following.		
Possible Clinical Effect	Probability of Occurrence	Treatment
Intracranial injections may exhibit temporary post-op clinical signs related to the procedure.	Post-op clinical signs occur infrequently following injection and typically resolve after a few days. We expect any clinical signs to be mild and not affect the animals' ability to locomote or eat.	Consult with veterinary staff if clinical signs develop

V. END POINT CRITERIA

A. What clinical signs will be used as a basis for removal of an animal from the study? If any animals begin to display neurological deficits or other clinical signs that may impact their health and wellbeing, including a Clinical Ratings Scale posture score of 3, gait score of 3 or more, bradykinesia score of 3 or more, or gross motor skills score of 3 or more in both arms, they will be referred to the veterinary staff for evaluation.

Weight loss in excess of 20% of ideal weight (as determined by veterinary staff based on body weight and body condition score) that does not resolve after two weeks of supportive treatment (as determined and provided in conjunction with the DACT veterinary team).

An animal that becomes laterally recumbent, or has difficulty locomoting or feeding themselves which does not resolve after one week of supportive treatment (as determined and provided in conjunction with the DACT veterinary team), or when determined by the DACT Veterinary Team to have reached an endpoint (veterinary discretion).

VI. EUTHANASIA

- A. List the primary method of euthanasia: Transcardial perfusion under anesthesia. If not perfusing, pentobarbital-containing euthanasia solution.
- B. If using a chemical or gas, complete the chart below: Various combinations of the following drugs may be used in coordination with euthanasia via injection of a euthanasia solution or perfusion.

Agent	Dose	Route	<u>Is this a DEA controlled</u> substance (Y/N)?	<u>Secondary method</u> used to confirm euthanasia
Pentobarbital-containing euthanasia solution	86-120 mg/kg	IV	Y	Removal of brain
Ketamine	10-20 mg/kg	IM, IV	Y	Used in conjunction with perfusion
Xylazine	2-4 mg/kg	IM, IV	N	Used in conjunction with perfusion
Midazolam	0.05-0.5 mg/kg	IM	Y	Used in conjunction with perfusion
Atropine	0.02-0.05 mg/kg	IM	N	Used in conjunction with perfusion
Morphine	1-2 mg/kg	IM, IV	Y	Used in conjunction with perfusion
Hydromorphone	0.2-0.4 mg/kg	IM, IV	Y	Used in conjunction with perfusion
Heparin	5,000 IU	IC	N	Used in conjunction with perfusion
Isoflurane	3-5%	Inhalation	N	Used in conjunction with perfusion
Sevoflurane	5-8%	Inhalation	N	Used in conjunction with perfusion
0.9% saline	1-2 L	IC	N	Used in conjunction with perfusion
4% formaldehyde	1-2 L	IC	N	Used in conjunction with perfusion

C. If euthanasia is being done by a physical means (e.g., decapitation, cervical dislocation) without anesthesia, provide scientific justification: N/A

I. BLOOD COLLECTION

A. Will blood be collected?

 \square No. Proceed to section II. \bowtie Yes. Complete the following

res. Complete the following.					
Site	Volume (ml)	% BW	Max. # of	Min. Interval	
		// BIT	collections		
femoral vein	≤10 mL	≤0.5%	Up to 5 planned,	Typically 4 weeks;	
			7 max including	Rarely within 7	
			potential redraws	days (see below)	

- B. Will anesthetics, sedatives, or other drugs be used during blood collection?
 - No. Proceed to section I. C.
 - Yes. Complete the following.

Drug	Dose	Route	Purpose
Ketamine	3-10 mg/kg	IM	Anesthesia
Dexmedetomidine	0.03 mg/kg	IM	Anesthesia
Midazolam	0.05-0.5 mg/kg	IM	Anesthesia
Atropine	0.02-0.05 mg/kg	IM	Reduce respiratory
			secretions

C. Describe the methods used to draw the blood including physical restraint, if any. Animals will be anesthetized with ketamine and either dexmedetomidine or midazolam. Blood samples will be obtained from the femoral vein and separated for serum collection. Sedation is expected to last 30 minutes.

D. Provide scientific justification for blood collection and justification for the frequency of it. Plasma and serum will be used for measuring inflammatory biomarkers. Collections spaced approximately a month apart will allow for adequate bodily replacement. In the event that a blood collection attempt does not yield an adequate sample (i.e.: no sample, inadequate volume [<1 mL plasma/serum]), the blood collection may be repeated up to one additional time within a 7-day period, well below accepted blood draw volume levels and frequency limits for animals of this size (2 – 10 kg).

II. OTHER TISSUE/BODY FLUID COLLECTION

- A. Will other tissues or body fluids be collected prior to death?
 - No. Appendix 1 is completed.
 - Yes. Complete the following. Surgical procedures should be described more fully in Appendix 2.

Tissue/Fluid	Site and Method	Amt	# of collections	Min Interval
CSF	lumbar or	≤0.5 mL	Up to 5 planned,	Typically 4 weeks;
	cisternal puncture		7 max including	Rarely within 7
			potential redraws	days (see below)

- B. Will anesthetics, sedatives, or other drugs be used during tissue/body fluid collection?
 - No. Proceed to section II. C.
 - Yes. Complete the following.

Drug	Dose	Route	Purpose
Ketamine	3-10 mg/kg	IM	Anesthesia
Dexmedetomidine	0.03 mg/kg	IM	Anesthesia
Midazolam	0.05-0.5 mg/kg	IM	Anesthesia
Atropine	0.02-0.05 mg/kg	IM	Reduce respiratory secretions
Betadine/Isopropyl alcohol	N/A	Topical	Topical disinfectant

Revised 2/25/2021 Obtained by Rise for Animals. Uploaded to Animal Research Laboratory Overview (ARLO) on 08/15/2023 C. Describe the methods used to collect the samples, including physical restraint, if any.

Animals will be anesthetized with ketamine and either dexmedetomidine or midazolam. CSF collection is performed as a sterile procedure. The lumbar or cervical area of the animal will be shaved and scrubbed alternating with povidone iodine and alcohol three times to prepare the collection site. A sterile drape will be placed over the collection site and sterile surgical gloves will be worn for the collection. For lumbar collection, a 22G spinal needle will be advanced into the spinal subarachnoid space until CSF begins to flow spontaneously. For cisternal collection, a 22G spinal needle attached to a 3-6 mL syringe will be advanced into the cisterna magna and CSF will be withdrawn. Sedation is expected to last 30 minutes.

D. Provide scientific justification for the sample collection(s) and justification for the frequency of it CSF will be used for measuring inflammatory biomarkers. Collections spaced approximately a month apart will allow for adequate bodily replacement. In the event that a CSF collection attempt does not yield an adequate sample (i.e.: no sample, inadequate volume [<0.2 mL], or blood contamination), the CSF collection may be repeated up to one additional time within a 7-day period.

APPENDIX 2: SURGICAL PROCEDURES

I. GENERAL INFORMATION

- A. Species Cynomolgus macaque
- B. Surgical Procedure(s) Intracranial injection
- C. Room/location of surgery Surgical Suite

II. PRE-SURGICAL CARE

- A. Will the animals undergo pre-surgical fasting?
 - No. Proceed to section III.

Yes. Provide the details:

The day before a scheduled surgical procedure, animals are offered their full diet allotment in the early afternoon, and any remaining diet is removed at the end of the workday. The animal is then fasted overnight until the scheduled surgery the following morning in order to mitigate the risk of emesis and aspiration during the procedure.

III. SURGICAL PROCEDURE:

Survival 🗌 Nonsurvival

*Note: A surgical checklist is recommended for each survival surgery, and possibly non-survival surgeries. These checklists should be submitted to DACT's Research Support Services for review before implementing procedures.

A. Describe each surgical procedure (e.g., approach, tissue manipulation, closure): Intracranial Injections:

Anesthesia will be induced with injectable anesthetics, and animals will then be intubated and maintained on gas anesthesia. Morphine or hydromorphone will be administered pre-operatively as will Cefazoin Ani s will be placed in stereotaxic frames. Surgical targeting will be accomplished using a surgical neuronavigation system, which will allow in-op visualization of the surgical instruments within and around the brain. The MRI images will be uploaded to the system and coordinates for target areas will be marked. Under sterile conditions an 8 cm incision will be made along the midsagittal plane of the scalp. Entry points will be identified using the system. One entry hole will be drilled on each hemisphere of the skull (10mm x 10mm). Animals will receive bilateral injections of rAAV-Cav1.3-shRNA (n=1) or rAAV-Scr-shRNA (n=2) in the putamen (3 sites per hemisphere, 15 µL in the rostral and middle sites and 10 µL in the caudal site). Infusion will syringes will be lowered be performed with an infusion pump attached to a stereotaxic micromanipulator to the targets, and the contents infused at a rate of 1 µL/min. After the injection is complete, the needle/syringe will be left in place for an additional 2 minutes to allow infusate to diffuse from the needle tip and prevent backflow prior to retracting the syringe. The entry holes will be filled with Gelfoam. The SC tissues, and skin will then be closed using absorbable suture. Bupivacaine (1-2 mg/kg, SQ) will be administered to the incision site prior to closure. Anesthesia depth will be monitored approximately every 10 minutes through jaw tone, palpebral reflex and vitals measurements (e.g., heart rate, ventilatory rate, oxygen saturation, blood pressure, temperature, CO2 level). Animals will receive Buprenorphine SR and Meloxicam. Surgery is expected to last approximately 2 hours. Should any animal experience adverse effects post-surgery (including signs of infection) as determined by the veterinary staff, they will be evaluated and treated as appropriate.

B. Anesthetic regimen:

Various combinations of the following drugs may be used in the induction and maintenance of anesthesia for surgery.

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Drug & concentration (e.g., mg/ml)	Dose (e.g., mg/kg) & maximum volume to be given	Route	Is this a DEA controlled substance (Y/N)?
Ketamine (100 mg/mL)	10-15 mg/kg, 1 mL	IM	Υ
Midazolam (5 mg/mL)	0.05-0.5 mg/kg	IM	Υ
Dexmedetomidine	0.015-0.05 mg/kg	IM	Ν
Atropine (0.54 mg/mL)	0.02-0.05 mg/kg	IM	Ν
Isoflurane	0.5-5%	Inhalation	Ν
Sevoflurane	1-8%	Inhalation	Ν
Propofol (10 mg/mL)	2-5 mg/kg, 5 mL (Bolus) 0.2-0.6 mg/kg/min (CRI)	IV	Ν
Cetacaine Spray (Benzocaine 14%, Butamben 2%, Tetracaine 2%)	200 mg	Topical	N

Please refer to the IACUC approved document "Macaque Anesthesia/Analgesia/Antibiotics Regimens"

- Note: Use of gas anesthetics requires completion of the EH&S-based Anesthetic Gas Safety training prior to use and refreshed annually.
- 1. Describe measures used to indicate a surgical plane of anesthesia to keep animals from getting too light or too deep:

Anesthesia depth will be monitored approximately every 10 minutes through jaw tone, palpebral reflex, and vitals measurements (e.g., ECG, heart rate, ventilatory rate, oxygen saturation, blood pressure, temperature, end tidal gases).

C. Additional pharmacological agents used during surgery (include analgesics, supportive medications, and research drugs):

Drug and concentration	Dose & max volume	Route	Purpose	Frequency	Is this a DEA controlled substance (Y/N)?
Betadine/Chlorhexidine/ Isopropyl alcohol	N/A	Topical	Topical Disinfectant	Once, as needed	N
Bupivacaine (5 mg/mL)	1-2 mg/kg, 2 mL	SC	Analgesia	Once during closure	N
Cefazolin (330 mg/mL)	20-25 mg/kg, 0.76 mL	IV	Antibiotic	Every 2-4 hours, intraoperatively	N
Gelfoam	Cut to size	Topical	Hemostasis/Seal surgical holes	Once, as needed	Ν
Hydromorphone (2 mg/mL)	0.05-0.2 mg/kg	SC, IM, IV	Analgesia	Once, as needed	Y
Morphine	1-2 mg/kg	IM, IV	Analgesia	Once, as needed	Y
Ophthalmic ointment	Dab	Topical	Prevent corneal desiccation	Once, as needed	N
Saline or Lactated Ringer's Solution	5-15 mL/kg/hr, 300 mL	IV	Fluid replacement	Constant-rate infusion	N
Sufentanil (0.5 µg/mL)	0.25-2 µg/kg/hr, 120 mL	IV	Analgesia	Constant-rate infusion	Y

D. Describe the steps taken to maintain an aseptic surgery: Trained individuals will perform standard sterile prep of the scalp. The site will be scrubbed alternating with

povidone iodine/chlorhexidine and alcohol three times. Sterile drapes, gowns, gloves, and instruments will be used.

E. What is the maximum duration of each surgery? 4 hours

F. Will any animals recover from surgery?

No. This involves terminal, or non-survival, procedures; Appendix 2 is complete.

 \boxtimes Yes. Complete Section IV.

IV. POST-SURGICAL CARE

- A. Is there a potential for post-operative pain or distress?
 ☐ No. Proceed to section C.
 ☑ Yes.
- B. Will analgesics be used?

(For analgesic options, refer to the IACUC Standard Institutional Guideline on analgesia (https://researchintegrity.asu.edu/animals/procedures-library-and-guidelines) or contact a DACT veterinarian

□ No. Provide a scientific justification:

\boxtimes	Yes.	Complete	the	following.
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Drug & concentration	Dose & max. volume	Route	Frequency	Is this a DEA controlled substance (Y/N)?
Buprenorphine Sustained release (1-3 mg/mL)	0.2 mg/kg	SC	Once post- op	Y
Meloxicam Sustained release (10 mg/mL)	0.6 mg/kg	SC	Once post- op	Ν
Meloxicam (5 mg/mL injection; 1.5 mg/mL oral)	0.1-0.2 mg/kg	SC, PO	SID as needed/ variable duration based on procedure	Ν

Please refer to the IACUC approved document "Macaque Anesthesia/Analgesia/Antibiotic Regimens"

Who will administer these drugs? Veterinary staff or other trained individuals.

- C. Post-operative routine care:
 - i. What other drugs will be administered, if any (e.g., antibiotics, fluids)?

Drug & concentration	Dose & max. volume	Route	Purpose	Frequency	<u>Is this a DEA</u> <u>controlled</u> substance (Y/N)?
Cefazolin (330 mg/mL)	20-25 mg/kg	IM	Antibiotic	BID as needed/ variable duration based on procedure	N
Cephalexin (50 mg/mL)	20-30 mg/kg	PO	Antibiotic	BID as needed/ variable duration based on procedure	N

Please refer to the IACUC approved document "Macaque Anesthesia/Analgesia/Antibiotic Regimens" (choice of antibiotic and route of administration dictated by patient compliance. We try oral administration first, but default to injectable if NHP is not compliant)

ii. What other post-operative support and monitoring will be provided, how often, for how long, and by whom? Pain assessment scoring is performed following major surgical procedures and continues until the pain score is 0 as determined by the veterinary or trained research staff. Monitoring is provided by both trained DACT and PI personnel. Should any animal experience adverse effects post-surgery (including signs of cerebral infection, cranial incision complications, or neurologic deficits) as determined by the veterinary staff, they will be evaluated and treated as appropriate by the veterinary staff.

D. Is post-operative intensive care required?

No. Proceed to section E.

What special care is required?

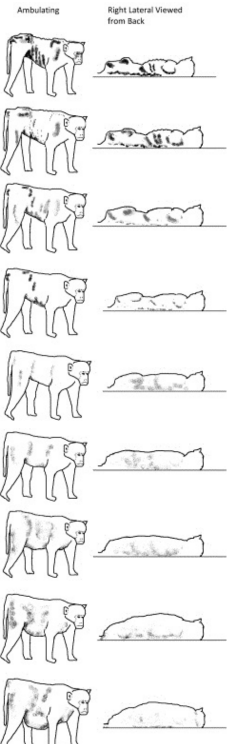
Who will provide special care and what are their qualifications?

For how long will special care be needed?

- E. Will animals undergo multiple survival surgical procedures?
 - \boxtimes No. Appendix 2 is complete.
 - Yes. Describe which surgeries, the sequence (specifying time between surgeries), and frequency. Provide scientific justification:

9-point Body Condition Score

1	EMACIATED – Very prominent hip bones (easily palpable and likely visible), prominent facial bones, spinous processes and ribs. Minimal to no muscle mass is palpable over ileum or ischium. Anus may be recessed between ischial callosities. Body is very angular, no subcutaneous fat layer to smooth out prominences.
2	VERY THIN – Hips, spinous processes, and ribs are prominent. Facial bones may be prominent. There is very little muscle present over the hips and back. Anus may be recessed between ischial callosities. Body is angular, no subcutaneous fat to smooth out prominences
3	THIN – Very minimal fat reserves, prominent hip bones and spinous processes. Hips, spinous processes and ribs are easily palpable with only a small amount of muscle mass over hips and lumbar region.
4	LEAN – Overlying muscle gives hips and spine a more firm feel. Hip bones and spinous processes are readily palpable, but not prominent. Body is less angular because there is a thin layer of subcutaneous fat.
5	OPTIMUM –Hip bones, ribs and spinous processes are palpable with gentle pressure but generally not visible. Well developed muscle mass and subcutaneous fat layer gives spine and hips smooth but firm feel. No abdominal, axillary or inguinal fat pads.
6	SLIGHTLY OVERWEIGHT – Hip bones and spinous processes palpable with firm pressure but are not visible. Bony prominences smooth. Rib contours are smooth and only palpable with firm pressure. Small abdominal fat pad may be present.
7	HEAVY – Bony contours are smooth and less well defined. Hip bones, spinous processes and ribs may be difficult to palpate due to more abundant subcutaneous fat layer. May have fat deposits starting to accumulate in the axillary, inguinal or abdominal areas.
8	OBESE – This animal will often have prominent fat pads in the inguinal, axillary or abdominal region. Abdomen will be pendulous when animal sitting or ambulating. Hip bones and spinous processes difficult to palpate. Bony contours smooth and poorly defined.
9	GROSSLY OBESE –Obvious, large fat deposits in the abdominal, inguinal and axillary regions. Abdominal palpation is very difficult due to large amount of mesenteric fat. Pronounced fat deposits may alter posture/ambulation. Hip bones, rib contours and spinous processes only palpable with deep palpation.



PRR22-11_0597

IACUC Protocol Trackable Components Checklist

Protocol #: 22-1886R If for amendm	ent, amendment #:
PI:	
Species: NHP	Highest Category of Pain: D
Completed by	Date completed: 10/22/21
No trackable components in th	is document
Exceptions to the Guide:	
to half the normal allotment to provide period will last up to one month, and p weeks. Food restricted animals will be	ng the hand reach task may have their biscuit allotment restricted e incentive for the task. Restriction during the initial task training post-surgery restriction may last up to one week out of every 4 weighed weekly, or twice per week if body mass loss exceeds 8% o or more body weight or whose BCS drops to 3.5/9 or below will
Prolonged Restraint Species: Details:	
	de table pairing partners are not available. Animals that will undergo exempt from establishment of new social housing pairs.
Other:	
Other Trackable Components:	
Survival Surgerie(s) Species: NHP Surgerie(s): Intracranial injectio Multiple Major?: Yes XI	
	d level): AAV constructs cant, toxin, irritant, carcinogen, etc.): 4% Formaldehyde UV light, lasers, noise, magnetic fields, etc.): MRI (magnetic fields

Non-Centralized Animal Housing

IACUC Protocol Trackable Components Checklist

Location: Maximum duration:

Decapitation

USDA-covered Species exempt from USDA reporting

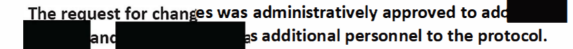
Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

	Animal	Protocol	Review
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ASU Protocol Number:	22-1886R RFC 1
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
	Antidyskinetic Therapy for PD
Principal Investigator:	
Date of Action:	12/6/2021

The animal protocol review was considered by the Committee and the following decisions were made:



NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training.or contact Research Support Services within DACT a t

Additional requirements:

Protocol Approval Period:

🗌 This protocol requires that Research Su	upport Services group wit	hin DACT provide supervision for the
first time a procedure is conducted. Conta	ac to so	chedule.
\square This protocol indicates that there are s	urgical procedures. A sur	rgical checklist may be required to be
submitted to Research Support Services w	vithin DACT	prior to starting surgeries.
⊠ Other requirements	nust be added to IBC disc	losure before working with
biohazardous materials.		

Total # of Animals:	3	
Species:	NHP	Pain Category: D

11/18/2021 - 11/17/2024

Sponsor: ASU Proposal/A Title:	Award #:	
Signature:	IACUC Chair or Designee	Date: 12/17/2021
Cc:	IACUC Office; IACUC Chair	



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name:	Phone:	
Dept: ASU-Banner Neurodegenerative	Email:	
Disease Research Center		

Participant #1	Add to: IBC # IACUC #21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1887R Delete from: IBC # IAC.UC #	FOR ORIA USE ONLY Training Verification
Name	ASURITE: Email:	
Project Responsibili	ties in IBC:	
Experience/Train	ing in These Responsibilities:	
which procedures a supervision: Intracr PET scan, blood/CS	re they responsible for on the IACUC protocol (please note re being done independently and which are done under anial surgery, intracarotid surgery, intracisternal injection, MRI, F collection, behavioral tests, administration of medications, and direct supervision until certified).	11/2021 OHSP
17 years' experience Experienced with int blood/CSF collection	Mice Experience and training with species and procedures: e in primate research. 14 years' experience in rodent research. racranial surgery, intracarotid surgery, MRI, PET scan, n, behavioral tests, administration of medications, and necropsy. racisternal injection by Dr.	

Participant #2	Add to: IBC #		
Name:	ASURITE:	Email:	
Project Responsibil		Linar	
Experience/Train	ing in These Responsi	ibilities:	
What procedures a	re they responsible fo	r on the IACUC protocol (pleas	e note 11/2021
which procedures a	re being done indepe	ndently and which are done u	nder OHSP
supervision: Intract	anial surgery, intracard	otid surgery, intracisternal injec	tion, MRI,
	F collection, behaviora direct supervision until	al tests, administration of medic certified).	ations, and
Species: Macaques,	Mice Experience and	training with species and proc	edures: 2
Experienced with int behavioral tests, add intracarotid surgery	racranial surgery, MŘl mi <u>nistration of m</u> edi <u>cat</u>	ears' experience in rodent rese I, PET scan, blood/CSF collecti ions, and necropsy. Will be trained will be trained rnal injection by Dr	on, ned in

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with animals or biological materials as applicable.

Principal Investigator Signature:

Date: 12/2/2021

Revised 11/20/12

Obtained by Rise for Animals. Uploaded to Animal Research Laboratory Overview (ARLO) on 08/15/2023

FOR ORIA USE ONLY	IBC Approved	IACUC Approved 12/6/2021
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Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal Protocol Review

ASU Protocol Number: Protocol Title: 22-1886R RFC 2 Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent <u>Antidyskinetic The</u>rapy for PD

Principal Investigator: Date of Action:

12/10/2021

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was administratively approved to adc as additional personnel to the protocol.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at

Additional requirements:

Ducto col Annuoval Deviado

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact the first time a procedure is conducted. Contact the first time a protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT prior to starting surgeries.
 Other requirements the added to IBC disclosure before working with biohazardous materials.

Total # of Animals:	3	
Species:	NHP	Pain Category: D

FlotocorAppro	vai renou.	11/10/202	.1-11/1//2	024	
Sponsor: ASU Proposal/ Title:	Award #:				
Signature:	IACUC Chair or	Designee			Date: 12/17/2021
Cc:	IACUC Office				

11/10/2021 11/17/2024



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name:	Phone:
Dept: ASU-Banner Neurodegenerative Disease Research Center	Email:

Participant #3	Add to: IBC # IACUC #21-1867 22-1873R, 22-1880R, 22-1880 Delete from: IBC # IACUC #	
Name:	ASURITE: Email:	
Project Responsibili	ities in IBC:	
Experience/Train	ning in These Responsibilities:	
which procedures a supervision: Intracr PET scan, blood/CS	re they responsible for on the IACUC protocol (plea re being done independently and which are done ranial surgery, intracarotid surgery, intracisternal inje F collection, behavioral tests, administration of med direct supervision until certified).	under & NHP ection, MRI, 12/2021
years' experience in Experienced with inf	Mice Experience and training with species and pro- primate research. 4 years' experience in rodent res- tracranial surgery, MRI, blood/CSF collection, behave dications, and necropsy. Will be trained in intracaro will be trained in PET scan by nal injection by Dr.	earch. <i>i</i> oral tests,

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with animals or biological materials as applicable.

Principal Investigator Signature:		Date: 12/2/2021
FOR ORIA USE ONLY	BC Approved	IACUC Approved 12/10/2021

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R RFC 3	
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent Antidyskinetic Therapy for PD	
Principal Investigator:		
Date of Action:	3/23/2022	

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was approved by Designated Review to add the use of mannitol during intracranial injection surgery.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact. Research Support Services within DACT at

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT prior to starting surgeries.
 Other requirements:

Total # of Anim Species:	als:	3 NHP	Pain Category: D	
Protocol Appro	val Period:	11/18/2021 - 11/17/20	024	
Sponsor: ASU Proposal/A Title:	Award #:			
Signature:	IACUC Chair ór	Designee		Date: 3/23/2022
Cc:	IACUC Office IACUC Chair			

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R RFC 2
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
	<u>Antidyskinetic The</u> rapy for PD
Principal Investigator:	
Date of Action:	3/23/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was approved by Designated Review to add the use of mannitol during intracranial injection surgery.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to contact the context of schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT prior to starting surgeries.
 Other requirements:

Total # of Anim Species:	als:	3 NHP	Pain Category: D	
Protocol Approval Period:		11/18/2021 - 11/17/20)24	
Sponsor: ASU Proposal/A Title:	Award #:			
Signature:	IACUC Chair or	Designee		Date: 3/23/2022
Cc:	IACUC Office IACUC Chair			

ARIZONA STATE UNIVERSITY

Institutional Animal Care and Use Committee

REQUEST FOR CHANGES TO AN APPROVED PROTOCOL

	22-1886R RFC 3	
Title:	Geneti <mark>c Silencing of Striatal</mark> CaV1.3 Calcium Channels as a P <u>uptent Antidyskinetic Therapy</u> for P)
Principal Investiga	tor: Email Address:	
If not PL whom sh	ould we contact for questions related to this amendment: Email Address:	

Funded Unfunded

Requested Change (check all that apply):

- New procedures to be performed complete Part A, and Appendix 1 and/or 2 as applicable, and sign assurance. New species and or an increase in the number of animals to be used complete Part A and sign assurance.
- New location of housing or procedures complete Part A and sign assurance.
- New personnel complete Part B and sign assurance.

Other (includes changes in dosages, funding, etc.) complete Part A and sign assurance.

A. Description of Requested Changes

For new procedures or additional animals that are USDA-covered species (all mammals EXCEPT mice and rats bred for research), list the Category of Pain:

For new procedures or additional animals that are not USDA-covered species, will there be the potential to involve more than slight or momentary pain or distress that will <u>NOT</u> be relieved with anesthetics, analgesics, tranquilizer drugs, or other methods for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia)? No Yes

If yes, describe and justify:

If you are adding a procedure that could create pain or distress, you need to include a literature search for alternatives. If you are adding a new survival surgery, submit a surgical checklist.

If you are requesting an increase in animal numbers, provide justification with supportive statistics.

If you are adding additional funding sources, provide the grant agency, grant title and ASU proposal or award number.

Describe the changes you are requesting. We would like to include the option to administer mannitol (1 g/kg, IV, up to 100 mL) via constant-rate infusion over 20 min alongside maintenance fluids during the intracranial injection surgery. Systemic administration of mannitol prior to vector injection has been shown to greatly improve AAV gene transfer in rats by increasing the number of cells transduced and vector distribution to the present of mannitol is within the range used clinically in humans to reduce intracranial pressure by increasing brain osmolarity, and the resultant hyperosmolarity should resolve within 24 hours

References:

B. Addition of Personnel

All personnel who work with animals are required to have animal care training within the last four years. ASU IACUC training modules can be completed at https://asu.col.qualtrics.com/ife/form/SV_b2b2XRXRs1309f. Personnel are required to have Level III training certification on file with the IACUC office in order to perform procedures independently (without supervision). Training can be received from DACT staff or PIs can have a member of their lab approved by DACT to be a Certified Trainer who can then train others. Simply being Level III certified does not allow the training of others. See the IACUC web site (https://researchintegrity.asu.edu/animals/training) for more information on training and Level III forms.

<u>* Procedures other than husbandry, handling, or behavioral testing</u> MUST be performed under supervision unless the person is Level III certified to conduct the procedure independently. Personnel are not Level III certified until the IACUC has reviewed and approved the Level III training documentation. The PI is responsible for ensuring that personnel who are not Level III certified are supervised at all times.

<u>Name Tit</u>	<u>ASURITE</u> name	What activities will each person perform on live animals ONLY while under direct supervision?	What activities will each person be allowed to perform independently (including appropriate Level 3 certification*) at the time of protocol submission?	Species with which individual will have direct contact ("all" or list species) *	IACUC USE ONLY Training (mm/yy)
		_			

For each individual, describe the individual's training and years of experience with all listed species and procedures they will be conducting under this protocol. For procedures for which they are not yet trained, but will likely be trained to do during the activity period of this protocol, provide a description of who will provide such training:

Assurance

As Principal Investigator of this protocol, I assure that all procedures will be conducted as described in this request for changes and that personnel will receive appropriate additional training prior to conducting any new procedures that are not listed above.

SIGNED	
Deineinel leventienten	<u>3/2/22</u>
Principal Investigator	Date
For IACUC use only: Administratively approved - Approving administ	
Administratively handled by VCV - Veterinarian Sources used for verification:	
Approved by Designated Review – Designated r Approved by Full Committee Review – Primary	

From: To: Subject: Date:



22-1886R RFC 3

I approve as the designated reviewer.

To: Subject:		acuc@asu.edu d: Designated Review for 22	1886R RFC 3
From Sent: Friday, Marc To: Cc: IACUC@asu.ed	h 18, 2022 11 22 AM Dale De	eNardo	Karen Kibler
Designated Revi Principal Invest Peer Reviewer: Protocol Numbe	igator: N/A	86R RFC 3	
Da	ale DeNardo aren Kibler	Response Yes: 3/18/2022 11:30 AM Yes: 3/23/2022 12:13 PM Yes: 3/18/2022 11:26 AM Yes: 3/18/2022 11:26 AM Yes: 3/18/2022 11:27 AM Yes: 3/18/2022 11:57 AM Yes: 3/18/2022 11:57 AM Yes: 3/18/2022 11:45 AM Yes: 3/18/2022 11:22 AM	

A request for an amendment to the referenced protocols has been submitted. The request for amendment is attached and the various related protocols are available at the SharePoint site. The Designated Review process allows IACUC members, by voting, to permit the Designated Reviewer to approve or disapprove an amendment request.

Select "YES" if you approve the use of the designated review process for this amendment.

Select "NO" if you disapprove the use of designated review process regarding this amendment request. The amendment will then be reviewed at the next monthly IACUC meeting.

Select "Abstain" if you would like to abstain from the vote for any reason.

Select "Recuse" if you have a conflict of interest.

Please indicate your approval or disapproval of the request for designated review by using the YES or NO button in the toolbar at the top of this message. The use of "YES or NO" buttons allows you to submit comments along with your choice. You may also send your comments to me directly or to the primary reviewer without using the selection buttons. Please copy me on all correspondence and email related to this request.

Sincerely,

Protocol #: 22-1886R If for amendment, amendment #: 3			
PI:			
Species: NHP Highest Cat	egory of Pain: D		
Completed by Date comp	leted: 3/17/22		
No trackable components in this document			
Exceptions to the Guide:			
to half the normal allotment to provide incentive for period will last up to one month, and post-surgery weeks. Food restricted animals will be weighed we	each task may have their biscuit allotment restricted r the task. Restriction during the initial task training restriction may last up to one week out of every 4 ekly, or twice per week if body mass loss exceeds 8% ly weight or whose BCS drops to 3.5/9 or below will		
Prolonged Restraint Species: Details:			
Husbandry Deviation from the Guide Species: NHP Deviation: Single housing if suitable pairing partners are not available. Animals that will undergo the hand reach task are permanently exempt from establishment of new social housing pairs.			
Other:			
Other Trackable Components:			
Survival Surgerie(s) Species: NHP Surgerie(s): Intracranial injection Multiple Major?: Yes No			
Hazardous Agents Biological (list agent and hazard level): AAV Chemical (note category toxicant, toxin, in Physical (note type - radiation, UV light, last and up to ~110 dB noise)			

Non-Centralized Animal Housing

Location: Maximum duration:

Decapitation

USDA-covered Species exempt from USDA reporting

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance Arizona State University

Animal Protocol Review

ASU Protocol Number: Protocol Title: 22-1886R RFC 4 Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent <u>Antidyskinetic The</u>rapy for PD

Principal Investigator: Date of Action:

4/21/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was approved by Designated Review to add the option for an additional craniotomy during intracranial injection surgeries.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT prior to starting surgeries.
 Other requirements:

Total # of Anim Species:	als:	3 NHP	Pain Category: D	
Protocol Appro	val Period:	11/18/2021 – 11/17/20)24	
Sponsor: ASU Proposal/A Title:	Award #:			
Signature:	ACUC Chair or	Designee		Date: 4/22/2022
Cc:	IACUC Office; I	ACUC Chair		

ARIZONA STATE UNIVERSITY

Institutional Animal Care and Use Committee

REQUEST FOR CHANGES TO AN APPROVED PROTOCOL

Protocol No. Title:	21-1867R, 22-1873R, 22-1880R, 22-1886R, 22-1887R, 22-1918R Differential diagnosis of Parkinson's and multiple system atrophy in non-human primate models using a novel a-synuclein retinal contrast agent and Al-assisted analytics
	Reprogramming astrocytes to functional dopaminergic neurons in non-human primate brain
	Co-Pathologies Drive Neuroinflammation and Progression in PD
	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent Antidyskinetic Therapy for PD
	MAW Trebalose in an NHP model of Alzheimer's DiseaseAAV-GBA Therapy in an NHP model of PD
Principal Investigator:	Email Address:
If not PI, whom should we	contact for questions related to this amendment: Email Address:

Funded Unfunded

Requested Change (check all that apply):

- New procedures to be performed complete Part A, and Appendix 1 and/or 2 as applicable, and sign assurance.
- New species and or an increase in the number of animals to be used complete Part A and sign assurance.
- New location of housing or procedures complete Part A and sign assurance.

New personnel – complete Part B and sign assurance.

Other (includes changes in dosages, funding, etc.) complete Part A and sign assurance.

A. Description of Requested Changes

- For new procedures or additional animals that are USDA-covered species (all mammals EXCEPT mice and rats bred for research), list the Category of Pain:
- For new procedures or additional animals that are not USDA-covered species, will there be the potential to involve more than slight or momentary pain or distress that will <u>NOT</u> be relieved with anesthetics, analgesics, tranquilizer drugs, or other methods for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia)? No Yes

If yes, describe and justify:

If you are adding a procedure that could create pain or distress, you need to include a **literature search** for alternatives. If you are adding a new survival surgery, submit a surgical checklist.

- If you are requesting an increase in animal numbers, provide justification with supportive statistics.
- If you are adding additional funding sources, provide the grant agency, grant title and ASU proposal or award number.

Describe the changes you are requesting. We would like to add the option to make one additional craniotomy to visualize the superior segittal sinus during intracranial injection surgeries under this protocol. Intraoperative navigation is generally highly accurate following initial skin registration (i.e., correlation of the MRI scan with with the the animal's actual position in the stereotaxic frame using fiducial marker locations or tracing the skin surface with a tracked instrument). However, it is occasionally necessary for the surgeon to confirm navigational accuracy after the skin has been retracted with an anatomical landmark that is clearly visible on MRL The superior sagittal sinus is ideal for this purpose and, prior to the adoption of intraoperative navigation with the visualizing the sinus was the primary method of establishing a mediolateral zero point for stereotaxic MRI coordinates in all surgical cases, as described by In cases where visualizing the sinus is deemed necessary, the surgeon will make a small craniotomy (up to 10 x 3 mm) along the mediolateral axis. It is usually not necessary for the craniotomy to fully penetrate the skull and the surgeon will stop once the sinus is visible through the bone. In the very rare occasions when the sinus is inadvertently penetrated, digital pressure with surgical gel foam is sufficient to control bleeding. The craniotomy will be filled with gel foam prior to wound closure, which will be as previously described.

References:

B. Addition of Personnel

All personnel who work with animals are required to have animal care training within the last four years. ASU IACUC training modules can be completed at https://asu.col.qualtrics.com/ife/form/SV b2b2XRXRs1309f. Personnel are required to have Level III training certification on file with the IACUC office in order to perform procedures independently (without supervision). Training can be received from DACT staff or PIs can have a member of their lab approved by DACT to be a Certified Trainer who can then train others. Simply being Level III certified does not allow the training of others. See the IACUC web site (https://researchintegrity.asu.edu/animals/training) for more information on training and Level III forms.

<u>* Procedures other than husbandry, handling, or behavioral testing</u> MUST be performed under supervision unless the person is Level III certified to conduct the procedure independently. Personnel are not Level III certified until the IACUC has reviewed and approved the Level III training documentation. The PI is responsible for ensuring that personnel who are not Level III certified are supervised at all times.

<u>Name</u> <u>Title</u>	<u>ASURITE</u> name	<u>What activities will each</u> person perform on live animals ONLY while under direct supervision?	each person be allowed to perform independently (including appropriate Level 3 certification*) at the time of protocol submission?	Species with which individual will have direct contact ("all" or list species) *	I <u>ACUC</u> USE ONLY Training (mm/yy)

For each individual, describe the individual's training and years of experience with all listed species and procedures they will be conducting under this protocol. For procedures for which they are not yet trained, but will likely be trained to do during the activity period of this protocol, provide a description of who will provide such training:

Assurance

As Principal Investigator of this protocol, I assure that all procedures will be conducted as described in this request for changes and that personnel will receive appropriate additional training prior to conducting any new procedures that are not listed above.

SIGNED	
Principal Investigator	<u>4/6/2022</u> Date
For IACUC use only: Administratively approved - Approving add Administratively handled by VCV - Vetering Sources used for verification: Approved by Designated Review – Designated Approved by Full Committee Review – Pring	arian providing verification: Date of verification: ated reviewer: Date of approval: 4/21/2022

From: To: Cc: Subject: Date: Attachments:	IACUC@asu.edu RE: Action Required: Designated Review for Tuesday, April 19, 2022 4:18:09 PM image002.png
Thanks	approve the modified amendment as the designated reviewer.
Good luck on th	e research
To Cc <iacucasu.edu< td=""><td>April 19, 2022 2:51 PM IACUC@asu.edu @mainex1.asu.edu></td></iacucasu.edu<>	April 19, 2022 2:51 PM IACUC@asu.edu @mainex1.asu.edu>
-	ion Required: Designated Review for Multiprotocol RFC
Laboratory Manage ASU-Banner Neuro	er degenerative Disease Research Center (NDRC)
From:	
Sent: Tuesday, To	April 19, 2022 2:34 PM
	<u>IACUC@asu.edu</u> @ <u>mainex1.asu.edu></u> ion Required: Designated Review for Multiprotocol RFC
	this outlines and justifies the process. This is very helpful. Please add a sentence nee to the amendment. Thanks!
From: Sent: Tuesday, J To	April 19, 2022 2:14 PM
	<u>IACUC@asu.edu</u> <u>@mainex1.asu.edu></u> ion Required: Designated Review for Multiprotocol RFC

Obtained by Rise for Animals. Uploaded to Animal Research Laboratory Overview (ARLO) on 08/15/2023 How the attached paper describes the stereotaxic surgery without the neuronavigation, including the exposure of the sagittal sinus for mediolateral zero. Is that what you are looking for?

Laboratory Manager

ASU-Banner Neurodegenerative Disease Research Center (NDRC)

From Sent: Tuesday, April 19, 2022 10:21 AM To Subject: FW: Action Required: Designated Review for Multiprotocol RFC

See below

Professor of Life Sciences

College of Liberal Arts and Sciences



Hi

From:	
Date: Tuesday, April 19, 2022 at 10:15 AM	-
То	
Cc	<u>c@asu.edu</u> " <u><iacuc@asu.edu></iacuc@asu.edu></u>
Subject: RE: Action Required: Designated Review for	Multiprotocol RFC

I am the designated reviewer for your amendment. I have no concerns, but can you please add some references to justify your discussion. Thank you!

Best,

Professor of Nutrition Arizona State University | College of Health Solutions



From
Sent: Tuesday, April 19, 2022 9:47 AM
То
Da <u>le DeNardo</u> Karen Kible
Cc: <u> ACUC@asu.edu</u>
Subject: Action Required: Designated Review for Multiprotocol RFC
Importance: High
Designated Reviewer:
Principal Investigator:
Peer Reviewer:
Protocol Number: Multiprotocol RFC

A request for an amendment to the referenced protocols has been submitted. The request for amendment is attached and the various related protoc s are available at the SharePoint site. The Designated Review process allows IACUC members, by voting, to permit the Designated Reviewer to approve or disapprove an amendment request.

Select "YES" if you approve the use of the designated review process for this amendment.

Select "NO" if you disapprove the use of designated review process regarding this amendment request. The amendment will then be reviewed at the next monthly IACUC meeting.

Select "Abstain" if you would like to abstain from the vote for any reason.

Select "Recuse" if you have a conflict of interest.

Please indicate your approval or disapproval of the request for designated review by using the YES or NO

To: Subject: RE: Action	iacuc@asu.edu Required: Designated Review for Multiprotocol RFC	
From Sent: Tuesday, April 19, 2022 1 To:	0:47 AM Dale DeNardo	Karen Kibler
Cc: IACUC@asu.edu Subject: Action Required: Desig Importance: High	nated Review for Multiprotocol RFC	
Designated Reviewer: Principal Investigator: Peer Reviewer: Protocol Number:	Multiprotocol RFC	
Tracking: Recipient Dale DeNardo Karen Kibler	Response Yes: 4/19/2022 3:25 PM Yes: 4/19/2022 4:18 PM Yes: 4/19/2022 11:56 AM Yes: 4/19/2022 11:02 AM Yes: 4/19/2022 10:49 AM Yes: 4/19/2022 12:20 PM Yes: 4/19/2022 4:08 PM Yes: 4/19/2022 12:53 PM Yes: 4/19/2022 11:51 AM	

A request for an amendment to the referenced protocols has been submitted. The request for amendment is attached and the various related protocols are available at the SharePoint site. The Designated Review process allows IACUC members, by voting, to permit the Designated Reviewer to approve or disapprove an amendment request.

Select "YES" if you approve the use of the designated review process for this amendment.

Select "NO" if you disapprove the use of designated review process regarding this amendment request. The amendment will then be reviewed at the next monthly IACUC meeting.

Select "Abstain" if you would like to abstain from the vote for any reason.

Select "Recuse" if you have a conflict of interest.

Please indicate your approval or disapproval of the request for designated review by using the YES or NO button in the toolbar at the top of this message. The use of "YES or NO" buttons allows you to submit comments along with your choice. You may also send your comments to me directly or to the primary reviewer without using the selection buttons. Please copy me on all correspondence and email related to this request.

1

Sincerely,

Protocol #: 22-1886R	If for amendment, amendment #: 4
PI	
Species: NHP	Highest Category of Pain: D
Completed by	Date completed: 4/13/2022
No trackable components in t	his document
Exceptions to the Guide:	
to half the normal allotment to provid period will last up to one month, and p weeks. Food restricted animals will be	ng the hand reach task may have their biscuit allotment restricted e incentive for the task. Restriction during the initial task training post-surgery restriction may last up to one week out of every 4 e weighed weekly, or twice per week if body mass loss exceeds 8% 6 or more body weight or whose BCS drops to 3.5/9 or below will
Prolonged Restraint Species: Details:	
	ide itable pairing partners are not available. Animals that will undergo exempt from establishment of new social housing pairs.
Uther:	
Other Trackable Components:	
Survival Surgerie(s) Species: NHP Surgerie(s): Intracranial injecti Multiple Major?: Yes X	
	rd level): AAV constructs icant, toxin, irritant, carcinogen, etc.): 4% Formaldehyde , UV light, lasers, noise, magnetic fields, etc.): MRI (magnetic fields

Non-Centralized Animal Housing

Location: Maximum duration:

Decapitation

USDA-covered Species exempt from USDA reporting

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal Protocol Review

ASU Protocol Number: Protocol Title: 22-1886R RFC 5 Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent <u>Antidyskinetic The</u>rapy for PD

Principal Investigator: Date of Action:

5/4/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was approved by Designated Review to revise the surgery details of the protocol.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT provide supervision for the prior to starting surgeries.
 Other requirements:

Total # of Anim Species:	als:	3 NHP	Pain Category: D	
Protocol Appro	val Period:	11/18/2021 - 11/17/20)24	
Sponsor: ASU Proposal/A Title:	ward #:			
Signature:	ACUC Chair or			Date: 5/4/2022
Cc:	IACUC Office; IA	CUC Chair		

ARIZONA STATE UNIVERSITY

Institutional Animal Care and Use Committee

REQUEST FOR CHANGES TO AN APPROVED PROTOCOL

Protocol No.	22-1886R RFC 5
Title:	Genetic Siloncing of Striatal CaV1.3 Calcium Channels as a Potent Antidyskinetic Therapy for PD
Principal Investig	ator: Email Address
16 and Olympic	e contact for questions related to this amendment: Email Address:
X Funded U	

Requested Change (check all that apply);

New procedures to be performed complete Part A, and Appendix 1 and/or 2 as applicable, and sign assurance. New species and or an increase in the number of animals to be used complete Part A and sign assurance.

New location of housing or procedures complete Part A and sign assurance.

New personnel complete Part B and sign assurance.

Other (includes changes in dosages, funding, etc.) complete Part A and sign assurance.

A. Description of Requested Changes

For new procedures or additional animals that are USDA-covered species (all mammals EXCEPT mice and rats bred for research), list the Category of Pain:

For new procedures or additional animals that are not USDA-covered species, will there be the potential to involve more than slight or momentary pain or distress that will <u>NOT</u> be relieved with anesthetics, analgesics, tranquilizer drugs, or other methods for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia)? No Yes

If yes, describe and justify:

If you are adding a procedure that could create pain or distress, you need to include a **literature search** for alternatives. If you are adding a new survival surgery, submit a surgical checklist.

If you are requesting an increase in animal numbers, provide justification with supportive statistics.

If you are adding additional funding sources, provide the grant agency, grant title and ASU proposal or award number.

Describe the changes you are requesting. We would like to correct the surgery description for the intracranial injection surgery. The previously approved description was copied from an earlier version of the study design and does not reflect to esurgeries performed on the other animals in this study i the final version of the study design as used in the surgeries is as follows. Changes from the previously approved description are highlighted.

Anesthesia will be induced with injectable anesthetics, and animals will then be intubated and maintained on gas anesthesia. Morphine or hydromorphone will be administered pre-operatively as mit Ce stereotaxic frames. Surgical targeting will be accomplished using the surgical instruments within and around the brain. The MRI images will be uploaded to the system and coordinates for target areas will be marked. Under sterile conditions, an 8 cm incision will be made along the midsagittal plane of the scalp. Entry points will be identified using the system, but if necessary a small craniotomy (up to 10 x 3 mm) along the mediolateral axis may also be made to visual the superior sagittal sinus to confirm navigational accuracy as detailed in RFC 4. One or more entry holes will be drilled on each hemisphere of the skull (up to 10mm x 10mm) dorsal to the injection sites. Animals will receive bilateral inj of rAAV-Cav1.3-shRNA (n=1) or rAAV-Scc-shRNA (n=2) in the putamen

Infusion will be performed with an infusion pump attached to a stereotaxic micromanipulator. Hamilton syringes will be lowered to the targets, and the contents infused at a rate of 1μ L/min. After each injection is complete, the needle/syringe will be left in place for an additional structure to allow infusate to diffuse from the needle tip and prevent backflow prior to retracting the syringe. The entry holes and any craniotomy sites will be filled with Gelfoam. The SC tissues, and skin will then be closed using absorbable suture. Bupivacaine (1-2 mg/kg, SQ) will be administered to the

Revised 2/25/2021 Obtained by Rise for Animals. Uploaded to Animal Research Laboratory Overview (ARLO) on 08/15/2023

PRR22-11_0623

incision site prior to closure. Anesthesia depth will be monitored approximately every 10 minutes through jaw tone, palpebral reflex and vitals measurements (e.g., heart rate, ventilatory rate, oxygen saturation, blood pressure, temperature, CO2 level). Animals will receive Buprenorphine SR and Meloxicam. Surgery is expected to last approximately

Should any animal experience adverse effects post-surgery (including signs of infection) as determined by the veterinary staff, they will be evaluated and treated as appropriate.

B. Addition of Personnel

All personnel who work with animals are required to have animal care training within the last four years. ASU IACUC training modules can be completed at https://asu.col.gualtrics.com/jfe/form/SV b2b2XRXRs1309f. Personnel are required to have Level III training certification on file with the IACUC office in order to perform procedures independently (without supervision). Training can be received from DACT staff or PIs can have a member of their lab approved by DACT to be a Certified Trainer who can then train others. Simply being Level III certified does not allow the training of others. See the IACUC web site (https://researchintegrity.asu.edu/animals/training) for more information on training and Level III forms.

<u>* Procedures other than husbandry, handling, or behavioral testing</u> MUST be performed under supervision unless the person is Level III certified to conduct the procedure independently. Personnel are not Level III certified until the IACUC has reviewed and approved the Level III training documentation. The PI is responsible for ensuring that personnel who are not Level III certified are supervised at all times.

<u>Name</u>	<u>Title</u>	<u>ASURITE</u> name	<u>What activities will each</u> person perform on live animals ONLY while under direct supervision?	What activities will each person be allowed to perform independently (including appropriate Level 3 certification*) at the time of protocol submission?	<u>Species with which</u> individual will have direct contact ("all" or list species) *	IACUC USE ONLY Training (mm/yy)

For each individual, describe the individual's training and years of experience with all listed species and procedures they will be conducting under this protocol. For procedures for which they are not yet trained, but will likely be trained to do during the activity period of this protocol, provide a description of who will provide such training:

Assurance

As Principal Investigator of this protocol, I assure that all procedures will be conducted as described in this request for changes and that personnel will receive appropriate additional training prior to conducting any new procedures that are not listed above.

SIGNED:		
		4/27/22
Principal Investigator		Date
For IACUC use only:		
Administratively approved - Approvin	g administrator: Date of a	ipproval:
Administratively handled by VCV - Ve Sources used for verification:	terinarian providing verification:	Date of verification:
Approved by Designated Review – De	•	Date of approval: 5/4/2022 of approval:

From: Sent: To: Subject:	Monday, May 2, 2022 10:43 AM iacuc@asu.edu Yes: Action Required: Designated Review for 22, 1886R RFC 5
Follow Up Flag: Flag Status:	Follow up Flagged

I approve as the designated reviewer.

Η

To: Subject:	RE: Action Requi	iacuc@asu.edu red: Designated Review for 22 1886R RFC 5	
	May 2, 2022 9:41 AM		
To	Dale	DeNardo	Karen Kibler
Subject: Action Importance: H	n Required: Designated	Review for 22-1886R RFC 5	
Designated I Principal Inv Peer Review Protocol Nur	vestigator: er: N/A	22-1886R RFC 5	
Tracking:	Recipient	Response	
		Yes: 5/2/2022 10:43 AM	
		Yes: 5/2/2022 9:47 AM	
	Dale DeNardo	Yes: 5/2/2022 11:47 AM	
		Yes: 5/2/2022 9:47 AM Yes: 5/2/2022 9:51 AM	
	Karen Kibler	Yes: 5/2/2022 9.51 AM Yes: 5/2/2022 10:01 AM	
		Yes: 5/2/2022 9:43 AM	
		Yes: 5/2/2022 12:49 PM	
		Yes: 5/2/2022 12:42 PM	

A request for an amendment to the referenced protocols has been submitted. The request for amendment is attached and the various related protocols are available at the SharePoint site. The Designated Review process allows IACUC members, by voting, to permit the Designated Reviewer to approve or disapprove an amendment request.

Select "YES" if you approve the use of the designated review process for this amendment.

Select "NO" if you disapprove the use of designated review process regarding this amendment request. The amendment will then be reviewed at the next monthly IACUC meeting.

Select "Abstain" if you would like to abstain from the vote for any reason.

Select "Recuse" if you have a conflict of interest.

Please indicate your approval or disapproval of the request for designated review by using the YES or NO button in the toolbar at the top of this message. The use of "YES or NO" buttons allows you to submit comments along with your choice. You may also send your comments to me directly or to the primary reviewer without using the selection buttons. Please copy me on all correspondence and email related to this request.

1

Sincerely,

Protocol #: 22-1886R	If for amendment, amendment #: 5
PI	
Species: NHP	Highest Category of Pain: D
Completed by	Date completed: 4/28/2022
No trackable components in th	nis document
Exceptions to the Guide:	
to half the normal allotment to provide period will last up to one month, and p weeks. Food restricted animals will be	ng the hand reach task may have their biscuit allotment restricted e incentive for the task. Restriction during the initial task training post-surgery restriction may last up to one week out of every 4 weighed weekly, or twice per week if body mass loss exceeds 8% o or more body weight or whose BCS drops to 3.5/9 or below will
Prolonged Restraint Species: Details:	
	de table pairing partners are not available. Animals that will undergo exempt from establishment of new social housing pairs.
Other:	
Other Trackable Components:	
Survival Surgerie(s) Species: NHP Surgerie(s): Intracranial injectio Multiple Major?: Yes X	
	d level): AAV constructs cant, toxin, irritant, carcinogen, etc.): 4% Formaldehyde UV light, lasers, noise, magnetic fields, etc.): MRI (magnetic fields

Non-Centralized Animal Housing

Location: Maximum duration:

Decapitation

USDA-covered Species exempt from USDA reporting

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R RFC 6
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
-	Antidyskinetic Therapy for PD
Principal Investigator:	
Date of Action:	7/6/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The reques	t for changes was administratively approved to a	dc
	as additional personnel to the protocol.	

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT a

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT to starting surgeries.
 Other requirements:

Total # of Anim Species:	als:	3 NHP	Pain Category:	D
Protocol Appro	val Period:	11/18/2021 – 11/17/2024		
Sponsor: ASU Proposal/A Title:	Award #:			
Signature:	IACUC Chair or	Designee		Date: 7/7/2022
Cc:	IACUC Office IACUC Chair			



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name	Phone	
Dept: ASU-Banner Neurodegenerative	Email:	
Disease Research Center		

Participant #1	Add to: □ IBC #SPROTO202100000070 □ IACUC # 21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R Delete from: □ IACUC # Delete from: □ IBC # □ IACUC #	FOR ORIA USE ONLY Training Verification
Name	ASURITE: Email:	
	ilities in IBC: Will handle AAV viral vectors, alpha-synuclein Human Lewy Body extracts, and mouse/rat/nonhuman primate ssue.	e Added in ERA
Experience/Traini macaque research	ng in These Responsibilities: 7 years' experience in rodent a with ASU DACT.	nd
What procedures	e 5/2019	
which procedures	OHSP	
supervision: Maca blood/CSF collection (all under direct su Rats: Intracranial se any medications, a Mice: Intracranial any medications, a	ppsy n of	
the second se	s, Rats, Mice Experience and training with species and	
procedures: 7 yea	rs' experience in rodent and macaque research with ASU DA	CT.

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with

animals or biological materials as applicable.

Principal Investigator Signature:		Date: 7/1/22
FOR ORIA USE ONLY	IBC Approved	IACUC Approved 7/6/2022

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal	Protocol	Review
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ASU Protocol Number:	22-1886R RFC 7
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent <u>Antidyskinetic The</u> rapy for PD
Principal Investigator:	
Date of Action:	8/2/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The reques <u>t fo</u>	or changes was administratively approved to ad c
and	as additional personnel to the protocol.

<u>NOTE:</u> If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at the second seco

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact the context of the context o

Total # of Anim Species:	als:	3 NHP	Pain Category:	D
Protocol Appro	val Period:	11/18/2021 – 11/17/2024		
Sponsor: ASU Proposal/A Title:	Award #:			
Signature:	IACUC Chair or	Designee		Date: 8/2/2022
Cc:	IACUC Office; IA	ACUC Chair		



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name:	Phon
Dept: ASU-Banner Neurodegenerative	Email
Disease Research Center	

Participant #1	Add to: IBC #SPROTO202100000070 IACUC # 21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R Delete from: IBC # IACUC #	FOR ORIA USE ONLY Training Verification
Name:	ASURITE: Email:	
Project Responsibil	ities in IBC: Will handle AAV viral vectors, alpha-synuclein	
preformed fibrils, Hu blood/CSF/brain tiss	uman Lewy Body extracts, and mouse/rat/nonhuman primate sue.	
Experience/Trainin	g in These Responsibilities: No previous experience	
What procedures a	re they responsible for on the IACUC protocol (please note	7/2022
which procedures a	are being done independently and which are done under	OHSP
supervision: Maca	ques: Intracranial surgery, intracarotid surgery, MRI, PET scan, n, behavioral tests, administration of medications, and necropsy	
	pervision until certified).	
Rats: Intracranial su	urgery, blood/CSF collection, behavioral tests, administration of	
	nd necropsy (all under direct supervision until certified).	
	surgery, blood/CSF collection, behavioral tests, administration of	
	nd necropsy (all under direct supervision until certified).	and see and see a second
•	, Rats, Mice Experience and training with species and	
procedures: No pre		
Participant #2	Add to: □ IBC #SPROTO202100000070 □ IACUC # 21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1880R, 22-1886R, 22-1887R, 22-1903R, 22-1918R 22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R □ IACUC # Delete from: □ IBC # □	FOR ORIA USE ONLY Training Verification
Name	ASURITE: Ernail	
Project Responsibil	ities in IBC: Will handle AAV viral vectors, alpha-synuclein	
	uman Lewy Body extracts, and mouse/rat/nonhuman primate	
Experience/Trainin	g in These Responsibilities: No previous experience	
What procedures a	re they responsible for on the IACUC protocol (please note	7/2022
•	are being done independently and which are done under	OHSP
•	ques: Intracranial surgery, intracarotid surgery, MRI, PET scan,	
blood/CSF collectio		
(all under direct sup		
Rats: Intracranial su	urgery, blood/CSF collection, behavioral tests, administration of	
	nd necropsy (all under direct supervision until certified).	
	surgery, blood/CSF collection, behavioral tests, administration of	
	nd necropsy (all under direct supervision until certified).	
•	, Rats, Mice Experience and training with species and	
procedures: No pre	vious experience	

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with animals or biological materials as applicable.

Principal Investigator Signature:		Date: 8/1/22
FOR ORIA USE ONLY	IBC Approved	IACUC Approved 8/2/2022

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R RFC 8
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
	Antidyskinetic Therapy for PD
Principal Investigator:	
Date of Action:	8/11/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes was approved by Designated Review to update the possible detrimental sequelae on the protocol.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT a

Additional requirements:

□ This protocol requires that Research Support Services grou	p within DACT provide supervision for the
first time a procedure is conducted. Contact	to schedule.
□ This protocol indicates that there are surgical procedures.	A surgical checklist may be required to be
submitted to Research Support Services within DACT	prior to starting surgeries.
Other requirements:	

Total # of A Species:	nimals:	3 NHP	Pain Category: D	
Protocol Ap	proval Period:	11/18/2021 – 11/17/202	4	
Sponsor: ASU Propos Title:	a l/Aw ard #:			
Signature:	IACUC Chair o	r <u>Designee</u>	Date: 8/11/2022	
Cc:	IACUC Office IACUC Chair			

ARIZONA STATE UNIVERSITY

Institutional Animal Care and Use Committee

REQUEST FOR CHANGES TO AN APPROVED PROTOCOL

Protocol No.	21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R
Title:	Differential diagnosis of Parkinson's and multiple system atrophy in non-human primate models using a novel a-synuclein retinal contrast agent and Al-assisted analytics
	Kinase activation in multiple system atrophy
	Reprogramming astrocytes to functional dopaminergic neurons in non-human primate brain
	Co-Pathologies Drive Neuroinflammation and Progression in PD
	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent Antidyskinetic Therapy for PD
	AAV Trehalose in an NHP model of Alzheimer's Disease
	BAG3 in Rodent Models of Neurodegenerative Disease
	Bifunctional intrabody targeting intracellular alpha-synuclein
	Primate Holding, Assessment, and Training
	AAV-GBA Therapy in an NHP model of PD
Principal Investigator:	Email Address:
	e contact for questions related to this amendment: Email Address:

Funded Unfunded

Requested Change (check all that apply):

New procedures to be performed – complete Part A, and Appendix 1 and/or 2 as applicable, and sign assurance.

New species and or an increase in the number of animals to be used – complete Part A and sign assurance.

New location of housing or procedures – complete Part A and sign assurance.

New personnel – complete Part B and sign assurance.

Other (includes changes in dosages, funding, etc.) – complete Part A and sign assurance.

A. Description of Requested Changes

For new procedures or additional animals that are USDA-covered species (all mammals EXCEPT mice and rats bred for research), list the **Category of Pain**:

For new procedures or additional animals that are not USDA-covered species, will there be the potential to involve more than slight or momentary pain or distress that will <u>NOT</u> be relieved with anesthetics, analgesics, tranquilizer drugs, or other methods for relieving pain or distress (e.g., negative conditioning, unrelieved post-surgical pain, death without euthanasia)? No Yes

If yes, describe and justify:

If you are adding a procedure that could create pain or distress, you need to include a **literature search** for alternatives. If you are adding a new survival surgery, submit a surgical checklist.

If you are requesting an increase in animal numbers, provide justification with supportive statistics.

If you are adding additional funding sources, provide the grant agency, grant title and ASU proposal or award number.

Describe the changes you are requesting. We would like to add additional possible detrimental sequelae.

Possible Clinical Effect	Probability of Occurrence	Treatment
Surgical and other procedures performed under anesthesia may rarely result in death or permanent disability due to hemorrhage, edema, thrombosis, infection, toxicity, or complications due to anesthesia.	Rare	Consult with veterinary staff if clinical signs develop. Euthanasia may be considered.

B. Addition of Personnel

All personnel who work with animals are required to have animal care training within the last four years. ASU IACUC training modules can be completed at <u>https://asu.co1.qualtrics.com/ife/form/SV_b2b2XRXRs1309f</u>. Personnel are required to have Level III training certification on file with the IACUC office in order to perform procedures independently (without supervision). Training can be received from DACT staff or PIs can have a member of their lab approved by DACT to be a Certified Trainer who can then train others. Simply being Level III certified does not allow the training of others. See the IACUC web site (<u>https://researchintegrity.asu.edu/animals/training</u>) for more information on training and Level III forms.

<u>* Procedures other than husbandry, handling, or behavioral testing</u> MUST be performed under supervision unless the person is Level III certified to conduct the procedure independently. Personnel are not Level III certified until the IACUC has reviewed and approved the Level III training documentation. The PI is responsible for ensuring that personnel who are not Level III certified are supervised at all times.

<u>Name</u>	<u>Title</u>	<u>ASURITE</u> name	What activities will each person perform on live animals ONLY while under. direct supervision?	What activities will each person be allowed to perform independently (including appropriate Level 3 certification*) at the time of protocol submission?	<u>Species with which</u> individual will have direct contact (<u>"all" or</u> list species) <u>*</u>	<u>IACUC</u> <u>USE ONLY</u> <u>Training</u> (mm/yy)

For each individual, describe the individual's training and years of experience with all listed species and procedures they will be conducting under this protocol. For procedures for which they are not yet trained, but will likely be trained to do during the activity period of this protocol, provide a description of who will provide such training:

Assurance

As Principal Investigator of this protocol, I assure that all procedures will be conducted as described in this request for changes and that personnel will receive appropriate additional training prior to conducting any new procedures that are not listed above.

SIGNED:

Principal livestigator	8/4 Dat	<u>/2022</u> e
For IACUC use only:		
Administratively approved - Approving administrator:	Date of appro	oval:
Administratively handled by VCV - Veterinarian providing	verification:	Date of verification:
Sources used for verification:		
	Anna Khina Da	to of approval: 0/44/2022

Approved by Designated Review – Designated reviewer: Karen K bler Date of approval: 8/11/2022 Approved by Full Committee Review – Primary reviewer: Date of approval:

From:	Karen Kibler
To:	
Cc:	IACUC@asu.edu
Subject:	Multiprotocol RFC
Date:	Tuesday, August 9, 2022 10:46:50 AM
Attachments:	Multiprotocol RFC 8.5.2022 Final.docx

Hella

The attached version is DR approved.

Thanks, Karen

To:		iacuc@asu.edu	
Subject:	RE: Action R		protocol RFC 8.5.2022
From			
Sent: Tuesda	y, August 9, 2022 9:	47 AM	
To:			
	I	Dale DeNardo	Karen Kibler
	an adu		
Cc: IACUC@a	on Required: Design	ated Review for Multiprotocol RFC 8.	F 2022
Importance:			5.2022
importance.	i iigii		
Designated	Reviewer:	Karen Kibler	
	westigator:		
Peer Review Protocol Nu		N/A Multiprotocol RFC 8.5.2022	
FIOLOCOLINI	linder.	Multiprotocor Krc 8.5.2022	
Tracking:	Recipient	Response	
_			
		Yes: 8/9/2022 10:15 AM	
		Yes: 8/9/2022 10:10 AM	
	Dale DeNardo	Yes: 8/9/2022 10:31 AM	
		Yes: 8/10/2022 7:56 AM	
	Karen Kibler	Yes: 8/9/2022 10:15 AM	
		Yes: 8/9/2022 11:17 AM	
		Yes: 8/9/2022 3:29 PM	
		Yes: 8/9/2022 10:13 AM Yes: 8/10/2022 5:06 PM	
		162. 0/10/2022 3:00 PW	

A request for an amendment to the referenced protocols has been submitted. The request for amendment is attached and the various related protocols are available at the SharePoint site. The Designated Review process allows IACUC members, by voting, to permit the Designated Reviewer to approve or disapprove an amendment request.

Select "YES" if you approve the use of the designated review process for this amendment.

Select "NO" if you disapprove the use of designated review process regarding this amendment request. The amendment will then be reviewed at the next monthly IACUC meeting.

Select "Abstain" if you would like to abstain from the vote for any reason.

Select "Recuse" if you have a conflict of interest.

Please indicate your approval or disapproval of the request for designated review by using the YES or NO button in the toolbar at the top of this message. The use of "YES or NO" buttons allows you to submit comments along with your choice. You may also send your comments to me directly or to the primary reviewer without using the selection buttons. Please copy me on all correspondence and email related to this request.

1

Sincerely,

Protocol #: 22-1886R	If for amendment, amendment #: 8	
PI		
Species: NHP	Highest Category of Pain: D	
Completed by	Date completed: 8/8/22	
No trackable components in the	nis document	
Exceptions to the Guide:		
to half the normal allotment to provid period will last up to one month, and p weeks. Food restricted animals will be	ng the hand reach task may have their biscuit allotment restricted e incentive for the task. Restriction during the initial task training post-surgery restriction may last up to one week out of every 4 e weighed weekly, or twice per week if body mass loss exceeds 8% 6 or more body weight or whose BCS drops to 3.5/9 or below will	
Prolonged Restraint Species: Details:		
the hand reach task are permanently e	ide itable pairing partners are not available. Animals that will undergo exempt from establishment of new social housing pairs.	
Other:		
Other Trackable Components:		
 Survival Surgerie(s) Species: NHP Surgerie(s): Intracranial injection Multiple Major?: Yes No 		
 Hazardous Agents Biological (list agent and hazard level): AAV constructs Chemical (note category toxicant, toxin, irritant, carcinogen, etc.): 4% Formaldehyde Physical (note type - radiation, UV light, lasers, noise, magnetic fields, etc.): MRI (magnetic fields and up to ~110 dB noise) 		

Non-Centralized Animal Housing

Location: Maximum duration:

Decapitation

USDA-covered Species exempt from USDA reporting

Institutional Animal Care and Use Committee (IACUC) Office of Research Integrity and Assurance

Arizona State University

Animal Protocol Review

ASU Protocol Number:	22-1886R RFC 9
Protocol Title:	Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent
	Antidyskinetic Therapy for PD
Principal Investigator:	
Date of Action:	9/2/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The reques	<u>t for ch</u> anges was administratively approved to add
	s additional personnel to the protocol.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at the second seco

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT prior to starting surgeries.
 Other requirements: IBC approval of new personnel is required before work with biohazardous materials may begin.

Total # of Anin Species:	nals:	3 NHP	Pain Categor y: D
Protocol Appro	oval Period:	11/18/2021 – 11/17/2024	
Sponsor: ASU Proposal/ Title:	Award #:		
Signature:	IACUC Chair or	Designee	Date: 9/7/2022
Cc:	IACUC Office; IA	CUC Chair	



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name:	Phone
Dept: ASU-Banner Neurodegenerative	Email:
Disease Research Center	

Participant #1	Add to: IBC #SPROTO202100000070 [21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1887R, 22-1898R, 22-1901R, 22-1903R, Delete from: IBC # IACUC #	22-1886R, Training Verification
Name	ASURITE: Email:	
Project Responsil preformed fibrils, blood/CSF/brain t		
Experience/Train	erience	
What procedures	ol (please note 8/2022	
which procedures supervision: Mac blood/CSF collect (all under direct su Rats: Intracranial any medications, Mice: Intracranial any medications,	, MRI, PET scan, ons, and necropsy administration of certified). , administration of	
Species: Macaque		
procedures: 1 yea		

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with animals or biological materials as applicable.

Principal Investigator Signature:		Date: 9/1/22
FOR ORIA USE ONLY	BC Approved	IACUC Approved 9/2/2022

Animal Protocol Review

ASU Protocol Number:	22-18
Protocol Title:	Gene

22-1886R RFC 10 Genetic Silencing of Striatal CaV1.3 Calcium Channels as a Potent Antidyskinetic Therapy for PD

Principal Investigator: Date of Action:

9/23/2022

The animal protocol review was considered by the Committee and the following decisions were made:

The request for changes	was administratively	approved to ad	d
and	a	s additional per	sonnel.

NOTE: If you have not already done so, documentation of Level III Training (i.e., procedure-specific training) will need to be provided to the IACUC office before participants can perform procedures without supervision. For more information on Level III requirements see https://researchintegrity.asu.edu/animals/training, or contact Research Support Services within DACT at the second seco

Additional requirements:

This protocol requires that Research Support Services group within DACT provide supervision for the first time a procedure is conducted. Contact to schedule.
 This protocol indicates that there are surgical procedures. A surgical checklist may be required to be submitted to Research Support Services within DACT to schedule.
 Other requirements: IBC approval of new personnel is required before work with biohazardous materials may begin.

Total # of Anim Species:	als:	3 NHP	Pain Categor y : D
Protocol Appro	val Period:	11/18/2021 – 11/17/2024	
Sponsor: ASU Proposal/A Title:	Award #:		
Signature:	IACUC Chair or IACUC Office; I		Date: 9/27/2022



PERSONNEL MODIFICATION FORM IACUC and IBC

This form can be used to add or remove participants from both an IACUC protocol and an IBC disclosure. Please submit only one form to <u>Research.Integrity@asu.edu</u> and it will be processed by both committees.

Principal Investigator Name:	Phon
Dept: ASU-Banner Neurodegenerative	Email
Disease Research Center	

Participant #1	Add to: IBC #SPROTO202100000070 IACUC # 21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R, 22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R Delete from: IBC # IACUC #	FOR ORIA USE ONLY Training Verification
Name:	ASURITE Email:	
	ities in IBC: Will handle AAV viral vectors, alpha-synuclein	Already on IBC
	Iman Lewy Body extracts, and mouse/rat/nonhuman primate	in ERA
blood/CSF/brain tiss		
Experience/Trainin	g in These Responsibilities: No previous experience	
What procedures a	re they responsible for on the IACUC protocol (please note	8/2022
•	are being done independently and which are done under	OHSP
•	ques: Intracranial surgery, intracarotid surgery, MRI, PET scan,	
•	n, behavioral tests, administration of medications, and necropsy	
	ervision until certified).	
Rats: Intracranial su	Irgery, blood/CSF collection, behavioral tests, administration of	
	d necropsy (all under direct supervision until certified).	
	urgery, blood/CSF collection, behavioral tests, administration of	
	d necropsy (all under direct supervision until certified).	
	Rats, Mice Experience and training with species and	
procedures: No pre	vious experience	
	Add to: 🛛 IBC #SPROTO202100000070 🖂 IACUC #	FOR ORIA USE ONLY
Darticinant #2	21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R,	Training Verification
Participant #2	22-1887R, 22- <u>18</u> 98R, 22-1901R, 22-1903R, 22-1918R	
	Delete from: BC # ACUC #	
Name	ASURITE	
Project Responsibil	ities in IBC: Will handle AAV viral vectors, alpha-synuclein	Being added in
preformed fibrils, Hu	uman Lewy Body extracts, and mouse/rat/nonhuman primate	ERA
blood/CSF/brain tiss	sue.	
Experience/Trainin	g in These Responsibilities: No previous experience	
What procedures a	re they responsible for on the IACUC protocol (please note	10/2018
which procedures a	are being done independently and which are done under	9/2022 NHP
supervision: Macao	ques: Intracranial surgery, intracarotid surgery, MRI, PET scan,	OHSP
blood/CSF collection		
(all under direct sup		
Rats: Intracranial su		
any medications, an		
	urgery, blood/CSF collection, behavioral tests, administration of	
	d necropsy (all under direct supervision until certified).	
	Rats, Mice Experience and training with species and	
procedures: 3 years	s experience working with mice in research	
	Add to: 🛛 IBC #SPROTO202100000070 🖂 IACUC #	FOR ORIA USE ONLY
Participant #3	21-1867R, 22-1872R, 22-1873R, 22-1880R, 22-1886R,	Training Verification
	22-1887R, 22-1898R, 22-1901R, 22-1903R, 22-1918R	

Delete from: IBC # IACUC #	
Name: ASURITE: Email:	
Project Responsibilities in IBC: Will handle AAV viral vectors, alpha-synuclein preformed fibrils, Human Lewy Body extracts, and mouse/rat/nonhuman primate blood/CSF/brain tissue.	Need to add in ERA
Experience/Training in These Responsibilities: No previous experience	
What procedures are they responsible for on the IACUC protocol (please note	9/2022
which procedures are being done independently and which are done under	OHSP
supervision: Macaques: Intracranial surgery, intracarotid surgery, MRI, PET scan, blood/CSF collection, behavioral tests, administration of medications, and necropsy (all under direct supervision until certified). Rats: Intracranial surgery, blood/CSF collection, behavioral tests, administration of any medications, and necropsy (all under direct supervision until certified). Mice: Intracranial surgery, blood/CSF collection, behavioral tests, administration of any medications, and necropsy (all under direct supervision until certified).	
Species: Macaques, Rats, Mice Experience and training with species and	
procedures: No previous experience	

Assurance

As Principal Investigator, I assure that personnel will receive appropriate training prior to working with

animals or biological materials as applicable.

 Principal Investigator Signature:
 Date: 8/25/22

 FOR ORIA USE ONLY
 IBC Approved
 IACUC Approved 9/23/2022