



ACD WORKING GROUP ON  
ENHANCING RIGOR,  
TRANSPARENCY, AND  
TRANSLATABILITY IN ANIMAL  
RESEARCH



FINAL REPORT

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## Membership

### Chairs

- Barbara Wold, Ph.D.  
Bren Professor of Molecular Biology  
California Institute of Technology
- Lawrence A. Tabak, D.D.S., Ph.D.  
Principal Deputy Director  
National Institutes of Health

### External Members

- Nancy Ator, Ph.D.  
Professor of Behavioral Biology  
Department of Psychiatry and Behavioral Sciences  
Johns Hopkins School of Medicine
- Laís Berro, Ph.D.  
Postdoctoral Fellow  
University of Mississippi Medical Center
- Eliza Bliss-Moreau, Ph.D.  
Associate Professor  
Department of Psychology  
Core Scientist  
California National Primate Research Center  
University of California, Davis
- Romer A. Gonzalez-Villalobos, M.D., Ph.D., FAHA  
Senior Principal Scientist, Cardiovascular & Metabolism Discovery  
Janssen Research & Development, LLC
- F. Claire Hankenson, D.V.M., M.S., DACLAM  
Attending Veterinarian  
Director, Campus Animal Resources  
Professor, Pathobiology and Diagnostic Investigation  
College of Veterinary Medicine  
Michigan State University
- Véronique Kiermer, Ph.D.  
Chief Scientific Officer  
PLOS
- Keisa Williams Mathis, Ph.D.  
Assistant Professor  
Department of Physiology  
University of North Texas Health Science Center
- Sarah Nusser, Ph.D.  
Vice President for Research  
Professor of Statistics  
Iowa State University
- Regina Nuzzo, Ph.D.  
Senior Advisor for Statistics  
Communication  
American Statistical Association
- Eric Prager, Ph.D.  
Associate Director, External Affairs  
Cohen Veterans Bioscience
- F. Daniel Ramirez, M.D., M.Sc.  
Cardiac Electrophysiology Fellow  
Centre Hospitalier Universitaire de Bordeaux  
IHU Liryc
- Karen Svenson, Ph.D.  
Senior Scientific Program Manager  
Research Scientist  
Jackson Laboratory

## USG Members

- Brian Berridge, D.V.M., Ph.D., DACVP  
Associate Director  
National Toxicology Program  
Scientific Director  
Division of the National Toxicology  
Program  
National Institute of Environmental Health  
Science  
National Institutes of Health
- Paul Brown, Ph.D.  
Associate Director for Pharmacology and  
Toxicology  
Office of New Drugs  
Center for Drug Evaluation and Research  
Food and Drug Administration
- Janine Clayton, M.D.  
Director  
Office of Research on Women's Health  
National Institutes of Health
- Joshua A. Gordon, M.D., Ph.D.  
Director  
National Institute of Mental Health  
National Institutes of Health
- Michael Lauer, M.D.  
Deputy Director for Extramural Research  
National Institutes of Health
- Robyn Lee-Stubbs, M.S., CPIA, PStat®  
Chair  
Statistician  
Institutional Animal Care and Use  
Committee  
United States Army Medical Research  
Institute of Chemical Defense
- Glenn Merlino, Ph.D.  
Scientific Director for Basic Research  
Center for Cancer Research  
National Cancer Institute  
National Institutes of Health
- Shai D. Silberberg, Ph.D.  
Director  
Office of Research Quality  
National Institute of Neurological  
Disorders and Stroke  
National Institutes of Health
- Carrie Wolinetz, Ph.D.  
Acting Chief of Staff  
Associate Director for Science Policy  
National Institutes of Health

## Executive Secretary

- Jordan Gladman, Ph.D.  
National Institutes of Health

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Barbara Wold, Ph.D.

Lawrence A. Tabak, D.D.S., Ph.D.

## Executive Summary

Animals are a precious and essential part of NIH-funded biomedical research. Our working group was charged with advising NIH how it can help researchers improve the rigor, transparency, and reproducibility of NIH research involving animals, paying close attention to the late stages of the translational pipeline that produce new treatments for human health and disease. The overarching goal is to allow all stakeholders to have full confidence in the quality and applicability of research findings from animal studies and to ensure that animal subjects are used with appropriate consideration of ethics and harm–benefit analysis.

Researchers, funders, publishers, and the public provide an impetus to increase rigor and transparency in animal research. In part, this push has been motivated by recent documented problems in replication and translatability of biomedical research. Yet positive driving forces exist, too, such as the need to keep pace with the rapid progress of science itself, which includes technological advances that allow today’s researchers to work with increasingly complex questions and small effect sizes. This progress demands concomitant advances in scientific research methods—the rigor of study design, the handling and analysis of data, and the reporting of results.

Investigators need NIH’s support and active participation to increase the rigor and transparency of their research. NIH must obtain and commit sufficient financial resources toward improvements and also effectively use incentives and oversight in the grant application, review, and funding process. It can also uniquely help investigators by identifying and promulgating best practices, investing in strengthening the animal research statistical workforce, and working to educate the scientific community and the public about ongoing challenges and achievements.

Our working group’s recommendations follow five themes: (1) improving study design and data analysis; (2) addressing incomplete reporting and questionable research practices; (3) improving selection, design, and relevance of animal models; (4) improving methodological documentation and results reporting; and (5) measuring and evaluating the costs and effectiveness of these efforts.

In Theme 1, we recommend that NIH help researchers improve their study design and data analysis by two means: enhancing the statistical training available to students and investigators and facilitating collaborations between animal researchers and statisticians. We also recommend that NIH find ways for experts to provide feedback on investigators’ study plans early in the research process, helping applicants share these plans by adding dedicated extra space to NIH research grant applications.

In Theme 2, we recommend that NIH address questionable research practices by raising awareness in the animal research community of the benefits of prospectively documenting study design and analysis plans. We also recommend that NIH launch pilot programs for prospective registration and registered reports to explore their feasibility and utility.

In Theme 3, we recommend that NIH help improve the selection, design, and use of animal models by establishing a framework in which investigators can explain the scientific relevance and rationale behind their chosen animal models. We also recommend that NIH ensure that animal researchers have appropriate venues in which they can exchange best practices for animal models. More broadly, we recommend that NIH fund research programs on comparative animal–human biology and provide

adequate research support for larger and long-lived species. Lastly, we recommend that NIH educate the public on the value of animal research for enhancing human health and reducing illness, and we recommend it also create a working group to explore non-animal modeling systems in biomedical research.

In Theme 4, we recommend that NIH help improve study methodology documentation by two means: ensuring that researchers know which factors related to animals' environment can affect research outcomes and thus are critical to document and providing support for researchers to document the long-term care of larger and long-lived animals. We also recommend that NIH improve complete reporting of results by setting expectations for the use of the ARRIVE 2.0 Essential 10 checklist and the inclusion of statistical measures of uncertainty and effect size.

In Theme 5, we recommend that NIH measure the costs and effectiveness of efforts to improve rigor, transparency, reproducibility, and translatability in animal research. Specifically, we recommend that NIH develop a program to assess the progress in implementing this report's recommendations; conduct and support analyses on elements of rigor and transparency in grant applications and publications; allow applicants to include budget justifications for efforts linked to enhancing rigor, transparency, and reproducibility; and work with scientists who demonstrate the highest levels of transparency and rigor to develop best practices.

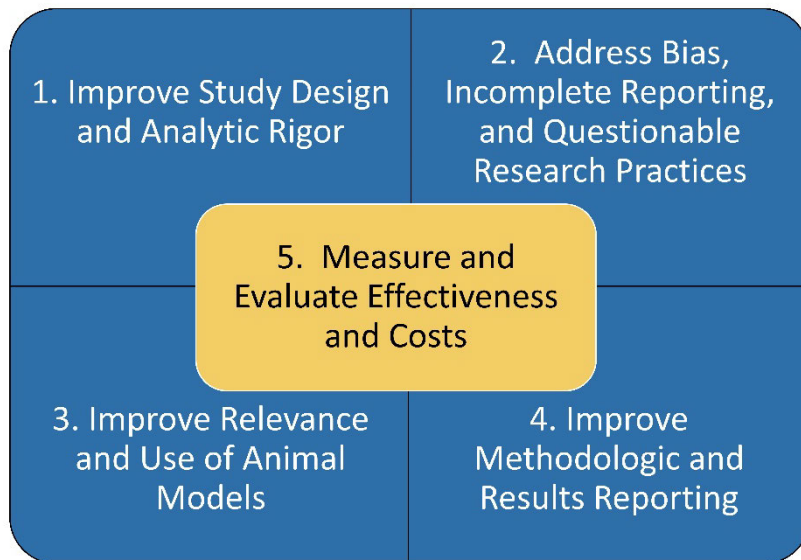


Figure 1. Themes of the ACD Working Group on Enhancing Rigor, Transparency, and Translatability in Animal Research Report

## Introduction

From understanding biological mechanisms to developing state-of-the-art vaccines, animal models play a critical role in science [\[1\]](#). Animal research has accounted for nearly half of all NIH research project grant applications over the past decade, and animal studies serve as a foundation for human clinical trials. Not only do we learn how to prevent, treat, and cure human diseases by studying animals, but often the treatments developed are also used to improve the health of animals themselves.

We study animals because of their comparative relationship to the human species. Moreover, unlike human studies, animal studies allow researchers to carefully control many characteristics that might affect the outcome of experiments, including intrinsic factors such as genetic composition and extrinsic factors such as diet or medications. Animal models thereby allow for a more precise understanding of biological factors and provide greater certainty about experimental outcomes.

Despite the numerous successes stemming from animal research, concerning reports over the past decade have described biomedical experiments that fail to replicate or to translate in ways that improve human health. All research is not expected to translate to human treatments, as there is no perfect model. Scientific process is as much about failure as it is about success. Yet part of the scientific process is also continual improvement, which includes working to understand what might contribute to unexpected outcomes within animal research.

In many fields of biomedical science, for example, animal researchers are now using more complicated statistical models to capture subtle effects, yet the statistical training and research support provided to investigators have not always kept pace. Likewise, although problems with questionable research practices and **publication bias** plague all researchers, solutions suitable for the animal research community are not in widespread use. Design and selection of animal models has become increasingly difficult, and it has been shown across multiple disciplines that extrinsic sources of variability from experimental conditions—such as lighting intensity, noise, vibrations and even the animal's microbiome—could contribute to difficulties in fully replicating experimental results.

The Institute for Laboratory Animal Research (ILAR) *Guide to the Care and Use of Animals in Research* [\[2\]](#) reflects, “Using animals in research is a privilege granted by society to the research community with the expectation that such use will provide either significant new knowledge or lead to improvement in human and/or animal well-being.” That expectation can be met only if studies are rigorous, well-designed, and transparently reported and use an animal model relevant to the condition of interest when intended for human **translation**. Following these practices will help ensure that the appropriate number of animals are involved to obtain reliable and reproducible results. Unreliable results from animal research can lead the scientific community astray, slow the progress of medical discovery, waste animals and other valuable resources, and lead the general public to lose trust in the scientific mission.

NIH has taken [several steps](#) in recent years to improve the scientific rigor of the research it supports, including releasing policies to enhance reproducibility of its supported research through rigor and transparency, increasing the focus of rigor in grant review, and establishing an [ACD working group](#) in response to the 21st Century Cures Act [\[3, 4\]](#). Yet animal research requires additional attention for several reasons. First, animal researchers are required to consider the ethical imperative of the “3 Rs”: **replacement, reduction, and refinement**. These principles impel researchers to use sound study design and contemporary methods for improved animal welfare and to include only the appropriate number of



animals necessary to answer a specific scientific question. Second, animal research faces unique challenges, such as species-specific animal husbandry and the appropriateness of selecting animal systems to model human biology and disease for translation. Third, even small improvements in animal research rigor can have a large impact on the funding of science, since animal studies serve as the foundational knowledge base for most human trials. Costs of research failures increase as research advances from early discovery-stage studies to mature translational animal studies that precede first-in-human trials. These costs include the misdirection of scientific effort, NIH dollars spent, and the well-being of human participants in follow-on patient trials. The resulting cost savings could then be reinvested in research at all stages from discovery to translation, thus further improving the return on investment.

Other organizations are also addressing reproducibility issues in animal research. In 2014, for example, the National Academies held a workshop on “The Missing ‘R’: Reproducibility in a Changing Research Landscape” [5], in which researchers discussed facets of animal-based research that could contribute to irreproducible results and provided perspectives on improving experimental planning, design, and execution; the importance of reporting all methodological details; and efforts to establish harmonization principles of reporting on the care and use of animals in research studies.

Improving rigor and reproducibility will have costs—in both dollars and opportunity—as well as benefits to science and society. Yet the amounts on both sides of the balance sheet are difficult to predict. Likewise, it will not be simple to adjust policies and practices in a data-driven way, but it is critical to do so. Managing the cost–benefit balance at the micro level (such as with individual research grants) and at the macro level (such as across the diverse NIH animal research portfolio) will be an ongoing challenge shared by NIH and its research community.

Rigorous science and its comprehensive reporting are primarily in the hands of researchers—in how we design, execute, and report our experiments, in how we conduct peer review of papers and grants, and in how we teach the next generation of scientists about rigor and transparency. The professional and societal benefits of conducting rigorous and transparent research are plentiful. Yet for individual animal researchers, the opportunity to elevate the 3 Rs can be a balancing act that often comes with obstacles in the form of added costs on “do-more-for-less” budgets, statistical training deficits, manpower constraints, misaligned incentives, and other challenges. NIH is uniquely positioned to support initiatives for improved scientific outcomes, building upon efforts it has already undertaken to improve rigor, transparency, and reproducibility, and taking into account work done by outside organizations, including the National Academies; the National Centre for the Replacement, Refinement, and Reduction of Animals in Research; the American Physiological Society; the Society for Neuroscience; the Federation of American Societies for Experimental Biology (FASEB); and other scientific societies.

## Working Group Charge

The Working Group was charged by the NIH Director to:

- Identify gaps and opportunities to improve the rigor, reproducibility, translational validity, and transparency of studies involving animal models, including:
  - Selecting animal models that are most appropriate for the scientific question

- Strengthening experimental design and statistical analysis, including appropriate statistical power and definition of endpoints
- Achieving appropriate transparency of methodological measures
- Considering how the conditions in which animals are housed and bred affect experimental outcomes
- Enhancing processes to incorporate rigor and transparency into grant applications, the peer review process, and manuscript publication
- Evaluate how animal models of human disease are currently developed, validated, and accepted into routine use and how this process could be improved
- Assess the current state of science for validating alternative models to animal research
- Consider the benefits and burdens of registering animal studies that aim to lead to first in human trials (e.g., preregistration of the experimental plan)
- Model the financial implications of potential changes in the average costs of grants using animal models, the number of studies funded, or the need to develop multi-lab consortia to achieve appropriate statistical power

## A Framework and Vocabulary to Discuss Rigor and Reproducibility

In our working group discussions, we quickly found that terminology and vocabulary played an important role in our shared understanding and framing. In this section and in [Appendix 1](#), we share the results from some of those discussions. While our charge did not specify a definition for “animal,” the major impetus for establishing the working group was to improve translation of animal research to human therapies. With this in mind, we focused our deliberations and emphasized recommendations that would most directly contribute to enhanced translatability. Although some recommendations and principles in the report could be applied to all biomedical research, our scope is “animals,” which we defined to include any non-human vertebrate or cephalopod species.

NIH defines “**scientific rigor**” as the strict application of the scientific method to ensure unbiased and well-controlled experimental design, methodology, analysis, interpretation, and reporting of results. Through the rigorous application of the scientific method, researchers can acquire knowledge that helps understand living systems and enhance health. **Transparency**, the accessibility of information, is essential for rigorous science and reproducibility.

“Reproducibility” and “replicability” are often used interchangeably to refer to sufficient consistency or robustness of experimental results when repeated by either the original researchers or, more often, by researchers who were not involved in the original experiments. However, these terms are also sometimes used to refer to distinct concepts. For example, we see both “reproducibility crisis” [\[6\]](#) and “replication crisis” [\[7\]](#) used to describe a growing concern among researchers, funding bodies, and the public that some of the scientific literature is fundamentally flawed or unreliable, and we read of a “replication project” [\[8\]](#) and “reproducibility projects” [\[9\]](#) designed to explore the prevalence of this problem. Despite the rapidly growing number of publications dealing with issues of “reproducibility” [\[10\]](#) and a consensus study by the National Academies of Sciences, Engineering, and Medicine [\[11\]](#), this term still lacks a standardized definition across the biomedical community. It was therefore necessary to clarify these terms and adopt a common vocabulary for the working group.

From the multiple definitions we reviewed, one set in particular resonated with our members. For our discussions and this report, we elected to follow terminology from the conceptual framework of reproducibility proposed by Goodman, Fanelli, and Ioannidis [\[10\]](#):

- **“Methods/methodological reproducibility”** refers to researchers providing enough detail about study procedures and data so that the same procedures could, in theory or in actuality, be exactly repeated. Transparency is a key component of methods reproducibility.
- **“Results reproducibility”** refers to researchers obtaining the same results when they conduct an independent new study with procedures as close to the original study as possible. This definition is similar to others’ definitions of “replicability.”
- **“Inferential reproducibility”** refers to researchers drawing qualitatively similar conclusions, or making knowledge claims of a similar strength, from either an independent replication of a study or a reanalysis of the original study. This is part of the process by which a scientific field decides which research claims or effects are to be accepted.

When assessing results reproducibility, it can be difficult for researchers to define what is required for results to be considered sufficiently similar. Some investigators use statistical significance (i.e., a  $p$ -value less than some threshold, typically 0.05) to determine whether a study has successfully replicated previous results. Yet this black-and-white approach to reproducibility can cause problems. For example, if a study captures an experimental effect with a  $p$ -value of 0.05, there is only a 50% chance that even a perfect replication study will also reach statistical significance [\[12\]](#) ([Appendix 2](#)). Expectations about “successful replications” should therefore be tempered. Perhaps more important than narrowly assessing replication is for researchers to assess the rigor of studies, consider the magnitude of their effects within the context of the field, and evaluate their cumulative evidence.

Similarly, when researchers evaluate inferential reproducibility, they must incorporate prior knowledge of experimental techniques and the subject matter at hand. If all results and analyses in a study are not completely reported, or if questionable research practices are suspected, then researchers should be less willing to accept the study’s research claim. In general, when undisclosed flexibility in study procedures increases, inferential reproducibility decreases. When experimental plans are documented and adhered to and study results are reported completely, inferential reproducibility increases.

Biomedical research is often conceptualized as spreading along a continuum: The earliest discovery research works to illuminate how biological systems work, and later **preclinical** research shifts to focus on knowledge to translate to human clinical trials. Human clinical trials often rely on applying results from late-stage preclinical animal studies; their success depends upon the quality of the preclinical research, so there is a cost when animal research fails to reproduce. Moreover, the costs stemming from low rigor and reproducibility grow as we move along the translational chain and radiate forward into therapeutic development in pharma and industry partners. Conversely, there are costs to insisting on high levels of rigor and reproducibility in early-discovery-stage research that may not be productive or beneficial. As part of our deliberations, we considered the unique needs and challenges of rigorous animal research across the stages of scientific pipeline.

The concept of **generalizability**, which an NSF report [\[13\]](#) defines as “whether the results of a study apply in other contexts of populations that differ from the original one,” is important to consider in the context of reproducibility and translation. In this report, we are concerned with whether results are

generalizable among animal studies and especially whether they can be generalized from an animal system to humans. We also recognize that there are different types of research using animals, which are generally designed and conducted in fundamentally different ways, and our methodological and inferential expectations of them should differ accordingly. Recognizing and embracing the distinction and co-dependence between **exploratory** (“hypothesis-generating”) and **confirmatory** (“hypothesis-testing”) research could also be beneficial in many ways. Journals, funding bodies, and peer reviewers should ensure that researchers who conduct exploratory experiments are not penalized for doing so. These researchers should feel encouraged to fully and transparently report experimental shortcomings (e.g., if experiments were underpowered or unblinded) and to appropriately temper the reach and strength of their conclusions rather than “sell” results as being more certain or being of more immediate translational potential than the evidence supports.

Meeting these unique challenges will require changes and investments from multiple angles. The itemized themes outlined below will highlight areas of opportunity for solutions and success.

## Theme 1: Improve Study Design and Data Analysis

Biomedical advances are largely founded on prior discoveries attained through experiments that are well-designed and carefully conducted and analyzed. When studies are rigorously designed in this way, they minimize the risk of unconscious researcher biases, control for random variability and confounders, and provide a reliable estimate of experimental effects. Key measures used to minimize the risk of bias include randomly and blindly assigning samples to comparison groups, conducting experiments and analyzing the results in a blinded fashion, and prospectively defining criteria for inclusion or exclusion of samples from analysis. Randomization serves to control for confounders by reducing the probability that comparison groups are not well-balanced. Prospectively defining a data analysis plan is also a fundamental aspect of good study design. Establishing a data analysis plan serves to refine the experimental design and ensures that questionable research practices associated with overly flexible analysis are mitigated (which we discuss in more detail in Theme 2).

The inherent variability among biological samples typically necessitates the use of multiple samples per group to achieve a reliable estimate of the parameters being measured. Optimal sample sizes minimize the risk of chance observations by reducing the probability that true relationships/effects will be overlooked or will lead to false-positive “findings.” Estimating an appropriate sample size is often complicated by insufficient information on the variability among the biological samples and expected effect sizes and by the complexity of the experimental design. Developing mature confirmatory designs requires antecedent and often stepwise exploratory work to refine these parameters. As such, it is imperative that scientists designing experiments or evaluating the work of others possess appropriate statistical skills and/or have access to statistical consultation.

Good study design and good data analyses are also essential from an ethical standpoint. Poorly designed studies can lead to mistaken results that lead the scientific community astray, slowing the progress of science. Such endeavors waste valuable resources and, importantly, might lead to unjustified use of animals or the need to repeat studies and waste animal life.

The following four recommendations under this theme address ways to help researchers improve their study designs, sample size estimations, and analysis plans and help reviewers and NIH assess the quality of such plans in manuscripts and grant applications.

**RECOMMENDATION 1.1: NIH SHOULD IMPROVE AND EXPAND STATISTICAL TRAINING FOR ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.1A: NIH SHOULD PARTNER WITH OTHER ORGANIZATIONS TO DEVELOP MODERN AND INNOVATIVE STATISTICS CURRICULA RELEVANT TO ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.1B: NIH SHOULD DEVELOP STATISTICAL RESOURCES SPECIFICALLY FOR ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.1C: NIH SHOULD REQUIRE STATISTICAL TRAINING FOR TRAINEES CONDUCTING ANIMAL RESEARCH AND STRONGLY ENCOURAGE IT FOR TEAM MEMBERS INVOLVED IN STUDY DESIGN AND DATA ANALYSIS.**

Animal researchers need strong statistical skills to produce research that is rigorous, reproducible, cost-effective, and ethical. To acquire these statistical skills, researchers need effective training. There are many dimensions to effectiveness in statistical education, but it is especially important that resources for animal researchers directly address the unique challenges and needs of doing research particularly with animal models. Training resources for animal researchers should also be easily accessible, engaging, and useful for audiences of different levels, from undergraduate students to principal investigators.

Current statistical training for animal researchers often falls short of these ideals, however. Curricula for research methods and statistics courses in graduate programs are not always modern, engaging, or relevant for animal researchers. Training requirements for NIH trainees vary greatly between programs. Resources for staying abreast of new statistical tools and methods for researchers beyond the trainee level are not easily accessible or centralized.

We recommend that NIH play a direct role in improving and expanding statistical training for animal researchers. A first step is to partner with other organizations (e.g., the American Statistical Association) to develop innovative and effective statistical curricula especially suitable for animal researchers. From this, specific resources should be developed and disseminated. The National Institute of Neurological Disorders and Stroke (NINDS) already has a planned educational platform on the principles of rigorous research; we recommend taking advantage of this opportunity and developing for this platform effective statistical resources specifically designed for animal researchers at various levels. Other platforms, such as the National Institute of General Medical Sciences (NIGMS) Clearinghouse for Training Modules to Enhance Data Reproducibility [\[14\]](#), can also be leveraged to enhance training. After these resources have been developed, we recommend requiring statistical training for NIH trainees conducting animal research, and we recommend strongly encouraging it for team members who are involved in study design and data analysis.

**RECOMMENDATION 1.2: NIH SHOULD FACILITATE COLLABORATION BETWEEN STATISTICIANS AND ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.2A: NIH SHOULD EXPAND RESEARCH COLLABORATIONS BETWEEN STATISTICIANS AND ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.2B: NIH SHOULD FUND TRAINING FOR STATISTICIANS ON DOMAIN-SPECIFIC SUBJECT MATTER AND ON CHALLENGES FACED BY ANIMAL RESEARCHERS.**

**SUB-RECOMMENDATION 1.2C: NIH SHOULD INCREASE ANIMAL RESEARCHERS' ACCESS TO STATISTICAL CONSULTING THROUGH FUNDING OPPORTUNITIES.**

**SUB-RECOMMENDATION 1.2D: NIH SHOULD INCENTIVIZE RESEARCH IN STATISTICAL METHODS FOR ANIMAL STUDY DESIGN AND ANALYSIS.**

Collaborations and cooperation between statisticians and empirical scientists are part of the culture in many scientific fields, including clinical research, social science, and agricultural science. This is not widespread practice in animal research, however. We recommend that NIH help change preclinical research culture by facilitating interactions between animal researchers and statisticians.

- *Research collaborations:* Collaborations between empirical scientists and statisticians can have a synergistic effect: Their interactions drive research advances in multiple areas, suggest new areas of research, and enhance both fields. We recommend that NIH find ways to facilitate research collaborations between animal researchers and statisticians through funded research programs. For example, institutions could establish short-term “idea incubator” residential educational programs that bring together junior statisticians and junior animal researchers where they are trained in issues at the interface of their fields, learn how to collaborate with each other, and build research relationships.
- *Training for statisticians:* When statisticians gain domain-specific scientific knowledge, they are able to communicate in the language of the scientific collaborator and better understand the underlying biological considerations in statistical approaches. We recommend that NIH fund training specifically designed for statisticians working with animal researchers. For example, training resources could be developed for graduate students on topics such as particular animal models, biological and scientific knowledge in specific fields, experimental design for animal studies, and applied statistics consulting skills.
- *Access to consulting:* When empirical scientists can consult applied statisticians, the additional methodological expertise can increase the rigor and effectiveness of their studies. We recommend that NIH increase animal researchers’ access to statistical consulting through funding opportunities. For example, seed funding could be provided for an institution to create a position specifically for an animal research statistical consultant or a faculty joint appointment between the Department of Statistics and the College of Veterinary Medicine. Additionally, graduate students in applied statistics could receive funding that would require statistical consulting projects with animal researchers.
- *Methodological research:* We recommend that NIH incentivize statistical methodology research specifically focused on animal study design and analysis. For example, research could be funded around small-sample techniques, methods for controlling for confounders common in animal experiments, strategies for randomization and blinding in animal experiments, and experimental designs and data analysis tools designed for specific animal models.

**RECOMMENDATION 1.3: NIH SHOULD ADD A SINGLE PAGE TO THE NIH GRANT APPLICATION RESEARCH STRATEGY SECTION THAT IS SOLELY DEDICATED TO THE DESCRIPTION OF CRITICAL ELEMENTS OF STUDY DESIGN, INCLUDING INCLUSION/EXCLUSION CRITERIA, SAMPLE SIZE ESTIMATION, DATA ANALYSIS PLAN, BLINDING, AND RANDOMIZATION, TO REDUCE THE RISK OF BIAS AND CHANCE OBSERVATIONS. THIS PAGE WOULD BE IN ADDITION TO THE CURRENT RESEARCH STRATEGY PAGE LIMIT AND WOULD APPLY TO VERTEBRATE AND CEPHALOPOD STUDIES.**

Journals and funders are increasingly requiring or encouraging researchers to report critical elements of their study design and analysis in reports and applications, often by using tools such as the ARRIVE and PREPARE guidelines and journal-specific checklists. Their goal is to increase transparency and reduce the

risk that published results are affected by undisclosed biases, random variability, or errors in experimental design, laboratory protocols, or data analysis. Unfortunately, investigators often fail to fully report all factors in their experimental design and analysis, and these journal efforts are continuing to evolve to improve and facilitate compliance [15, 16]. However, a major limitation of a journal's checklist is that it comes at the end of a study when it is too late to improve the design with a blinding and randomization plan or good sample size estimation.

Both the quality and character of future studies and the ease of using reporting checklists at the time of publication could be improved with two changes in approach by NIH. First, when investigators are fully aware of reporting recommendations and standards early in the research process, such as during the grant application process, they will naturally incorporate better design elements from the beginning. In this direction, a recent report from the Center for Scientific Review (CSR) Advisory Council [17] argues for increased consideration of rigor and reproducibility in all research in grant applications. Our working group concurs, and we identified additional needs for animal studies.

Second, an important lesson from studies of checklist successes and failures is that when fewer but more important items are recommended or required, investigators are more likely to fully report. Long reporting lists create burdens for investigators and hamper the ability of reviewers and editors to adequately evaluate or enforce compliance. We therefore concluded that the best path forward will be to include in the grant application Research Plan a succinct set of rigor items that are specifically tailored for animal use and that have already been found to improve reporting of blinding, randomization, and sample size calculations for *in vivo* studies.

Leveraging the recommendations from the CSR Advisory Council and acknowledging that short lists have increased value, we recommend that NIH have grant applicants who are doing vertebrate and cephalopod research explicitly address a specific list of study design elements as part of the Research Strategy. We also recommend that NIH provide supporting information to fully explain what is expected in the additional page (Appendix 3).

We recommend that these study design elements include:

- (1) *Inclusion and exclusion criteria*: criteria that will be used to include or exclude animals during data collection and analysis
- (2) *Sample size estimation*: how planned sample sizes for each group will be derived
- (3) *Data analysis plan*: what statistical tests will be used and how outliers will be defined
- (4) *Blinding*: who will be aware of group allocation at different stages of the experiment
- (5) *Randomization*: how allocation of experimental units to control and treatment groups will be done

We feel that the additional specificity required to satisfactorily address these elements of study design requires an additional, separate, dedicated page in the grant application. This additional page would be used solely for describing methods supporting specifically named elements and not for expanding upon or adding any other details of the Research Strategy. Applicants would use this page to summarize design strategies and further direct reviewers to sections in the Research Strategy where additional detail is provided. If any of these elements is not appropriate to the proposed studies, researchers would use this space to say that and explain why.



This dedicated single page would have several benefits. Most importantly, it would provide opportunities for researchers to sufficiently explain how these study design elements will improve the success of their study, which can subsequently be reported at the publication stages [15]. Given the multiplicity and diversity of study designs typically included in grant applications, this additional page would allow the investigator to give sufficiently detailed descriptions of bias-reducing measures that may not be permitted by the current space limitations. Furthermore, in support of the CSR Advisory Council and in line with Recommendation 1.4, reviewers at all stages of the process will be able to assess these elements and determine whether they have been adequately addressed and how they should affect funding or oversight. It would also enable NIH to retrospectively monitor the effectiveness of these measures across grants. Finally, this additional space would help level the playing field for young investigators or those new to later-stage preclinical work, many of whom do not have the option, often exercised by established investigators, of parsimoniously referring to rigor elements in a body of their own published work.

**RECOMMENDATION 1.4: NIH SHOULD EVALUATE WHERE IN THE PRE-STUDY RESEARCH PROCESS EXPERTS COULD ASSESS THE QUALITY OF STUDY DESIGN AND DATA PLANS, THEN IMPLEMENT PILOT STUDIES OF ASSESSMENT AT THE MOST PLAUSIBLE STAGE(S).**

A system of guardrails and quality checks in the research process can help ensure that researchers' study designs, data analysis plans, and statistical results are as strong and accurate as possible. We recognize the pressing need to have these "statistical guardrails" as well as the equally pressing obligation not to create an untenable burden for researchers, institutions, and NIH. Therefore, we recommend that NIH evaluate where in the pre-study research process statistical experts could assess the quality of study design and data plans, including the stages listed below, in order to balance enhanced rigor with acceptable burden. Expert assessment at multiple points in the research process might be appropriate. NIH should launch pilot studies to evaluate the effectiveness and assess the burdens of using quality checkpoints. Before implementation, a plan should be created to collect data and evaluate the efficacy and impact of the interventions.

- (i) *Researcher study design stage:* Grant applications with animal studies could be strongly encouraged to use the NC3R's Experimental Design Assistant tool [18, 19] and include its flowchart in the vertebrate animal section of the application.
- (ii) *Grant peer review stage:* All study panels with animal studies could include at least one trained reviewer who can evaluate the application's statistical elements, including study design and analysis plans. This would be aided by Recommendation 1.3, to create a separate page containing elements of study design and data analytic plans
- (iii) *Grant post-peer review stage:* As an alternative to (ii), a statistical review panel composed of applied statisticians can be formed to evaluate proposals with animal studies that have received the highest scores in the previous peer review stage. This would also be aided by Recommendation 1.3.

## Theme 2: Address Incomplete Reporting and Questionable Research Practices

The reliability of published research findings depends on the use of rigorous scientific practices designed to minimize unconscious research bias. Unfortunately, the use of "questionable research practices" has



been documented in many fields, including the absence of bias-controlling measures in experimental design [20-22], small and underpowered sample sizes, selective outcome reporting and outcome switching, *p*-hacking, and hypothesizing after the results are known (HARKing). These questionable research practices all increase the likelihood that study results and conclusions are unreliable.

Likewise, the reliability of published studies depends on the full reporting of research processes and outcomes. For example, when there is a lack of transparency in reporting how a study was conducted and analyzed or in reporting whether key parameters and procedures were decided before or after the data were collected, other researchers cannot fully interpret the study's results and evaluate its robustness [23, 24]. Similarly, when positive results receive priority for publication (a documented problem known as publication bias), the evidence base used to inform preclinical and clinical studies is weakened [25-27].

Strong motivation therefore exists to improve research and reporting practices, including the testing and adoption of approaches in which animal researchers plan and commit to experimental and analytic steps before they collect experimental data. The two recommendations under this theme address ways NIH can improve research and publication practices by addressing the problems of incomplete reporting and questionable research practices.

**RECOMMENDATION 2.1: NIH SHOULD LAUNCH A CAMPAIGN TO RAISE AWARENESS AND UNDERSTANDING OF PROSPECTIVELY DOCUMENTING STUDY DESIGN AND ANALYSIS PLANS.**

One approach to improving research rigor and transparency is prospective registration, in which researchers specify the details of their study design, analytic plan, and primary outcomes before data are collected and either register these details publicly (in a repository) or keep a time-stamped record under embargo until the study results are known. Prospective registration offers a way of retrospectively assessing bias control in study design by comparing published results with original plans. A publication process known as Registered Reports enables a journal to peer-review the study protocol before data collection and to offer a guarantee of publication regardless of the study results. In principle, this process offers benefits to researchers, helps improve study design before data collection, and protects against publication bias.

**Prospective registration:** Creating a permanent record of a study design, an analytic plan, and a primary outcome before the data are collected. Prospective registration allows retrospective insurance against selective reporting and outcome switching. The registered research plan can be embargoed for a limited time to protect intellectual property (IP), and when the registration is published, it allows identification and mitigation of publication bias. Researchers who prospectively register their studies demonstrate their intent to prioritize rigorous reporting. Prospectively registered studies do not undergo peer review at the time of registration.

**Registered Reports:** A journal article type in which the detailed study protocol is submitted for peer review before the data are collected. Upon successful review, the journal guarantees publication of the article regardless of the study findings. Like prospective registration, Registered Reports mitigate selective reporting and outcome switching. In addition, peer review of the protocol can help improve experimental design, for example, by ensuring that sample size and inclusion/exclusion criteria are predetermined and that proper measures are taken to mitigate experimental and analysis bias.

Researchers who publish with this review workflow demonstrate their intent to prioritize rigor, and they secure a path to publication regardless of findings, which mitigates publication bias.

Prospective registration has been successfully adopted in a number of research areas, including clinical research, economics [28], psychology [29], and systematic reviews in health and social care [30]. Despite implementation challenges, prospective registration has improved the accountability of clinical trials and transparency in their reporting [31-33]. Yet prospective registration is not in widespread use across all research disciplines, particularly in animal studies. Furthermore, awareness and understanding about what prospective registration means and its assumed benefits for research reliability are largely lacking in the animal research community. This lack of adoption and understanding in turn limits our ability to assess the efficacy of prospective registration in animal studies, demonstrate its benefits, measure its costs, and understand and mitigate its potential unintended consequences.

We recommend that NIH initiate a program to raise awareness and understanding of prospective registration methods and how they support universal goals of rigor and transparency. Key elements to communicate through this program include:

- Clear definitions of and distinction between prospective public (or embargoed) registration in a repository and Registered Reports, since these two mechanisms of prospective registration are likely candidates for potential pilot and intervention studies
- Clear articulation of the benefits of prospective registration and Registered Reports for researchers and their fields
- Mitigation measures (e.g., embargo periods) to be considered for legitimate concerns such as the protection of intellectual property and the minimization of risks to researchers, including if ongoing studies are discoverable by anti-animal research groups that engage in threatening behavior
- Awareness of prospective registration as a means to bring benefit to hypothesis-testing studies, particularly focusing on *in vivo* studies that are intended to directly inform clinical trials (i.e., toward the clinical end of the preclinical spectrum), as this is the context in which prospective registration can be most easily understood, justified, and applied

The campaign should begin with a strong message from NIH leadership—for example, in the form of a published commentary—to help raise the stakes and articulate the benefits sought through prospective registration.

**RECOMMENDATION 2.2: NIH SHOULD DEVELOP AND IMPLEMENT A PILOT PROGRAM TO GENERATE DATA ON AND EVALUATE THE EFFECTS OF SOLUTIONS THAT INVOLVE THE PROSPECTIVE DOCUMENTATION OF STUDY DESIGN AND ANALYSIS PLANS IN PRECLINICAL ANIMAL STUDIES.**

**SUB-RECOMMENDATION 2.2A: NIH SHOULD DEVELOP AND INCENTIVIZE PROJECTS THAT GENERATE DATA ON THE IMPACT OF PROSPECTIVE REGISTRATION AND REGISTERED REPORTS.**

**SUB-RECOMMENDATION 2.2B: NIH SHOULD SET UP A DEDICATED PROGRAM TO EVALUATE THE DATA GENERATED FROM THE PROJECTS RECOMMENDED IN 2.2A AND GUIDE FUTURE ADOPTION OF PROSPECTIVE REGISTRATION PRACTICES IN PRECLINICAL ANIMAL STUDIES.**

Because adoption of prospective registration in animal studies has been limited thus far, much remains unknown about its specific benefits, its cost to individual researchers and the research enterprise, and

its potential unexpected consequences. A path to broad adoption will require a stronger evidence base. However, based on the current lack of both understanding and appetite in the community, we believe that specific incentives will be required to generate sufficient data. We therefore recommend that NIH:

- Identify funding vehicles to incentivize prospective registration and Registered Reports for appropriate studies within a funded research proposal
- Develop an evaluation program to study the potential impact of these interventions and formulate a recommendation on future adoption of prospective registration

### **Prospective Registration Projects**

In order to gather data regarding the feasibility and utility of prospective registration to improve rigor and transparency in animal research specifically, we recommend that NIH support a prospective registration platform (possibly by partnering with existing registries) to allow any NIH grantee and applicant to prospectively register a research and analysis plan. If prospective registration were required, it would apply to hypothesis-testing studies that are typically a subset of studies described in a multi-year grant or are central to translational projects. Thus, to gather as much data as possible during an initial trial, we specifically recommend the following:

- NIH leadership should issue a call to action to all NIH grantees, incentivizing grantees to use the platform for prospective registration of studies within preexisting grants (with the option to embargo public release).
- Funding Opportunity Announcements (FOAs) specifically targeted at translational projects should serve as a testing ground for another prospective registration program. For a subset of such grants that have one or more Aims in which the proposed therapeutic is tested in animal models, NIH could require prospective registration of these specific studies. Prospective registration could be required as the first milestone in the study and described in a “prospective registration plan” to be submitted with the grant application; the following milestone would be the association of the prospective registration with the publication of the study results.

### **Registered Reports Projects**

The Registered Report publication format not only directly addresses publication bias but also presents strong benefits for researchers in the form of peer review of the proposed study design and a guaranteed publication regardless of study outcome. We recommend that this mode of publication be considered, where relevant, in NIH’s efforts to generate data. For example, NIH could encourage researchers to consider the Registered Report mode of publication, and it could incentivize its use by allowing evidence of an accepted “in-principle” Registered Report (i.e., post-protocol peer review but before publication of results) to be cited as an outcome in progress reports and grants and to influence reviewing and funding deliberations. Linking published Registered Reports to their grant IDs would allow the evaluation program to retrospectively evaluate how effective this publication format can be for reducing bias in the conduct and publication of studies.

### **Evaluation Program**

Before initiating a vehicle for gathering data on outcomes associated with prospective registration, we recommend that NIH establish a dedicated standing committee or task force to:

- Establish goals for collecting evidence related to prospective registration
- Specify or create funding programs that provide both a vehicle for data collection and approaches to analyzing outcomes
- Track the status of prospective registration plans submitted through annual NIH Research Performance Progress Reports (RPPRs)
- Publish a report of the outcomes of prospective registration projects and Registered Reports projects, specifying potential benefits and burdens of prospective registration at the NIH level, identifying funding programs for future prospective registration projects (assuming the data justifies further action), and clearly stating the future directions of the NIH prospective registration program

### Theme 3: Improve Selection, Design, and Relevance of Animal Models

Evolutionary conservation of mammalian biology is a fundamental feature supporting the wide use of animals as models for normal and diseased human biology. Animal models enable researchers to study biological and pathobiological processes at scales and under conditions untenable in humans. Our ability to recapitulate humanlike functional and morphologic phenotypes provides significant confidence in their human relevance. Yet human biology is complex, integrated, and individually variable. Because animal biology does not perfectly recapitulate humans, many animal models therefore imperfectly represent likely human outcomes.

A relatively small number of animal species have become staples in biomedical research representing varying degrees of human genetic, physiologic, and behavioral homology. Non-human primates generally represent the pinnacle of human-relatedness, though their use is minimized due to significant ethical sensitivities. Rodents, particularly mice, are the mammals most commonly used in biomedical research due to their small size, lower cost, and suitability for genetic manipulation. Other animal models, including zebrafish and invertebrates such as fruit flies and nematode worms, have also been useful for elucidating basic physiological processes.

Despite many similarities, differences in molecular, cellular, and organ systems–level physiology exist across species. Improving our approaches to animal model design and selection will require a multidisciplinary approach involving comparative animal scientists (representing animal biology), basic biomedical researchers (representing the questions and mechanism), and clinicians (representing the human context and outcomes of interest). When the scientific goal is direct translation to humans, an animal model should be characterized to ensure that fundamental elements of human biology are reasonably represented and that modulation would reflect a human response. Animal model selection should be rationalized by an evidence-based description of the human relevance of the model within the context of the experimental intent [\[34, 35\]](#).

High rates of failure in the development of novel therapeutics whose progression to human clinical trials was supported by animal studies has prompted concern that animal models of the human condition are more imperfect than presumed [\[36-38\]](#). That concern has instigated a useful and critical reflection on how and why we use animals in research as models either for humans or for particular aspects of human biology or behavior. Design and selection of animal models is one area of consideration, and we have made the following recommendations to strengthen the process.

**RECOMMENDATION 3.1: NIH SHOULD ESTABLISH A FRAMEWORK FOR RATIONALIZING THE SCIENTIFIC AND, WHEN APPROPRIATE, TRANSLATIONAL (HUMAN) RELEVANCE OF AN ANIMAL MODEL AND ITS SELECTION. THIS FRAMEWORK SHOULD BE EMPLOYED AS PART OF THE JUSTIFICATION FOR ANIMAL USES IN GRANT APPLICATIONS AND INCLUDED IN ETHICAL REVIEW PROCESSES AND IN JOURNAL REPORTS.**

Designing or choosing an animal model is not a straightforward undertaking. For many disorders, mechanisms of human disease cannot easily be translated [34, 39, 40]. In other cases, non-human species may not possess the biological features that are the target of modeling. Yet there is no established, standardized approach for selecting an animal model, and there is considerable variability in the approaches used to justify the use of a particular model in grant applications, journal articles, and other fora.

We recommend that NIH establish a framework for researchers to use to explain and justify the scientific relevance of their animal model and, when appropriate, its translatability to human processes and/or diseases. Such a framework would let applicants effectively justify their selected animal models and explain why the animal is appropriate for the specific scientific questions of interest. A standard framework would also help reviewers to evaluate the relevance of the animal models, which is especially important when the research seeks to help create new therapies, medical procedures, or diagnostics. With such a framework, reviewers would be better able to evaluate the scientific and public health impact of grant applications.

A useful framework could include a “points to consider” document or a question-based analysis that is provided with the supporting guidance for grant applications. It could include prompts to provide a brief summary of the known human biology or pathobiology, how that biology is represented in the animal, and historical experience with extrapolating outcomes from the model to humans. A few authors have proposed evidence-based frameworks that could guide these efforts [41-44].

**RECOMMENDATION 3.2: NIH SHOULD ESTABLISH OR IDENTIFY VENUES FOR THE EXCHANGE OF INFORMATION RELATED TO ANIMAL MODEL DESIGN AND CHARACTERIZATION, STUDY DESIGN, AND GENERAL BEST PRACTICES.**

While there are numerous avenues supporting and encouraging explication and exploration within specific scientific fields of inquiry or disease areas, there is little intellectual space devoted to understanding and promulgating the principles of proper design of animal models *per se*. The biomedical research community could benefit from dedicated pathways and processes for information exchange, training, and data collation and curation specifically related to the design and characterization of animal models. These various activities would be aimed at generating, maintaining, and disseminating a knowledge base focused on issues of rigor, reproducibility, and translatability in animal models. These activities would facilitate the exchange of data and experimental approaches that would promote the evaluation of disease models. Suitable venues could include journals, websites, or data platforms and might involve precompetitive consortia with private partners. The NCATS-funded Clinical and Translational Science Awards provide an existing infrastructure that could be leveraged.

**RECOMMENDATION 3.3: NIH SHOULD WORK TO IMPROVE THE DESIGN OF ANIMAL MODELS THROUGH THE FUNDING OF FOCUSED RESEARCH PROGRAMS THAT ENHANCE UNDERSTANDING OF COMPARATIVE HUMAN–ANIMAL BIOLOGY.**

A valuable target of support by NIH would be the funding of specific, cross-disciplinary scientific efforts that aim to develop and compare animal models for their utility in answering scientific and translational questions. These efforts would aid many fields by ensuring that comprehensive and unbiased processes

for evaluating models exist. These efforts to develop, characterize, and validate animal models of disease would also provide useful support and resource to disease-focused researchers. Spread across multiple laboratories and enabling collaboration across disciplines, these programs could also enhance reproducibility and enable effective and efficient training of the next generation of scientists. One mechanism for these efforts could be to employ centers of excellence (COEs) charged with providing leadership, best practices, research, support, and/or training with a specific focus area. Phenotypic and molecular characterizations as well as the outcomes of model validation studies could be shared in public resources such as those suggested in Recommendation 3.2. Continuing education courses in animal model design and use could be developed and provided.

**RECOMMENDATION 3.4: NIH SHOULD PROVIDE ADEQUATE RESEARCH SUPPORT FOR LARGER AND LONG-LIVED NON-RODENT SPECIES WHEN JUSTIFIED.**

**SUB-RECOMMENDATION 3.4A: NIH SHOULD CREATE POLICY TO ACCOMMODATE LONGER TIME FRAMES AND HIGHER BUDGETS FOR LARGER AND LONG-LIVED NON-RODENT SPECIES.**

**SUB-RECOMMENDATION 3.4B: NIH SHOULD CONTINUE TO DEVELOP NATIONAL RESOURCES TO PRODUCE LARGER AND LONG-LIVED ANIMALS.**

If grant funding needs to remain within the R01 budgets of \$500,000 (the limit per year without traversing an additional approval process), then the cost of acquiring and supporting larger and long-lived animals, particularly non-human primates, will limit longitudinal studies and will limit the ability to conduct experiments with sample sizes that provide sufficient statistical power. In studies of aging, for example, a 5-year funding period is insufficient for measurement of aging-related changes or pathological consequences of aging-related disorders. Therefore, NIH should provide both funding strategies and grant durations related to the species and experimental question being studied in larger and long-lived animals.

Often, discussions regarding the use of animals in research include a tacit argument that the use of small animals (or those with short lifespans) will save financial resources. In this view, the cost of each experiment or grant is the primary unit of measurement, and the lesser cost of smaller animals is evident. An alternative framework is to consider the costs to answer a particular research question or human health problem. In this view, a single large non-human primate study may cost substantially more than a mouse study, but if it can more effectively answer the research question and solve the human health challenge, it represents a better investment of resources.

The need to train young investigators in the care and use of larger and long-lived animals is crucial to using such species in translational or preclinical studies, but this is also resource intensive. Particularly important is training in how to choose the animal model that is most pertinent to the human disorder or disease of concern rather than vice versa. For example, this issue has been considered in relation to cardiology research [\[45\]](#) as well as for selecting optimal species for efficacy assessment in drug development [\[42, 44\]](#).

Gaps in funding are particularly problematic for research with larger and longer-lived animals. It is expensive to maintain such colonies, difficult to restart them, and almost impossible to start one *de novo* in the absence of a significant infusion of funds and institutional support (e.g., space, human resources support). Therefore, policies that ensure that facilities for such animals can maintain their colonies and retain staff with the relevant expertise across any funding hiatus are needed. Strategic

investments in existing resources (e.g., the National Primate Research Centers [46]) and/or generating new models of support for both large centers and smaller laboratories/programs are needed. Stagnant budgets to create and support national non-human primate breeding programs, for example, have significantly handicapped such centers, leading, in some cases (including COVID-19), to national shortages of experimental animals.

**RECOMMENDATION 3.5: NIH SHOULD EDUCATE THE PUBLIC ON THE VALUE OF ANIMAL RESEARCH, INCLUDING THE IMPORTANT ROLES OF LONG-LIVED, NON-RODENT MAMMALS FOR TRANSLATION TO IMPROVED HUMAN HEALTH AND DISEASE.**

Animals are essential in many areas and have produced medical advances that enhance the lives of both humans and the animal species originally studied. Animal research allows scientists to study animals throughout the entire life cycle and across generations within a manageable time frame. Laboratory conditions allow researchers precise control over the animals' environment and control experimental variables. Additionally, research with larger and long-lived animals, including non-human primates, has enabled the discovery and development of treatments and interventions for a variety of diseases. These research models have been instrumental to significant scientific and medical advances, including (but not limited to) deep brain stimulation to treat Parkinson's disease and experimental vaccines to prevent Ebola, polio, and COVID-19. NIH should take the strongest stances possible in supporting the need for research that uses animals properly by taking steps to educate the public on the value of this research regarding its translational relevance to cures for human disease. An additional focus should be on explaining the specific value of larger and long-lived animals.

There is also a need to improve the awareness and dissemination of existing resources and research networks provided to scientists using different species and working in different research areas to maximize translatability of research findings, especially for work with larger and long-lived animals. Initiatives such as the sharing of unpublished large animal data (e.g., "An Open Resource for Non-human Primate Optogenetics") and creation of large databases (e.g., the National NHP DNA Bank) should be expanded and promoted. Also, current initiatives for optimizing animal and tissue sharing need to be expanded and promoted across institutions and beyond the National Primate Research Centers.

**RECOMMENDATION 3.6: NIH SHOULD CHARTER A HIGH-LEVEL WORKING GROUP ON NON-ANIMAL MODELING SYSTEMS IN BIOMEDICAL RESEARCH TO COMPLEMENT THE ACTIVITIES AND RECOMMENDATIONS OF THIS ACD WORKING GROUP.**

Human cell-based modeling has also been a staple of biomedical research for more than 50 years, in which two-dimensional *in vitro* cultures of immortalized human cells have been the norm. These systems have enabled important investigation of human biology and disease at the cellular level without the need for animal studies, but they also have significant limitations depending on the question being asked. However, the rapid progress in our ability to model more complex elements of human biology in human-derived and *in vivo*-relevant modeling systems (e.g., human-derived microphysiological systems, organoids) is newly enabling study at the more complex tissue and organ levels [47-49]. Although still reductionist in overall biological complexity, these modeling systems enable more efficient (e.g., in time or cost) mechanistic investigation that could precede and complement animal studies or even replace them, depending on the intent of the research. Alternatives to animal models may be considered when a mechanistic question is amenable to a less complex modeling platform or when a translationally relevant animal model is lacking. The working group we envision would ideally be constituted by experts with experience in both developing and applying computational, *in vitro*, and *in vivo* modeling systems.



The group could examine similar questions regarding rigor, reproducibility, transparency, and relevance in the use of these systems, which likely share some considerations with animal studies but also come with their own sources of variance and uncertainty.

## Theme 4: Improve Methodological Documentation and Results Reporting

To achieve results reproducibility, methodological reproducibility must first be achieved. Researchers must provide enough details about the study details and procedures that independent investigators can repeat the study exactly. Methodological documentation and standards (e.g., ARRIVE [\[50, 51\]](#), CONSORT [\[52-56\]](#), MDRA [\[16\]](#), PREPARE [\[57\]](#)) for reporting results have been developed by the research community; these standards emphasize reporting critical elements in study design, sample size, inclusion and exclusion criteria, randomization, blinding, outcome measures, statistical methods, experiential animals, experiential procedures, and results. However, these standards are not currently enforced or required earlier on in the life of the study (e.g., grant applications, study design). Strengthening these elements across the life of a study, from planning to execution and publication, will result in a higher-quality knowledge base and will better inform future research.

**RECOMMENDATION 4.1: NIH SHOULD EXPECT THAT KEY SUPPORTING DATA REPORTED ON ANIMAL RESEARCH SUBMITTED IN SUPPORT OF GRANT APPLICATIONS WILL INCLUDE MEASURES OF QUALITY AND UNCERTAINTY FOR REPORTED ESTIMATES AND AN INTERPRETATION OF EFFECT SIZES WITHIN THE CONTEXT OF THE FIELD.**

For statistical results to be understood and properly interpreted, quantitative estimates must be accompanied by a measure of their quality or uncertainty, such as the standard deviation, confidence interval, credible interval, or posterior distribution of the cited estimate. Quantitative estimates are also most useful to other researchers when they are placed in the proper context, for example, by using standardized effect sizes or interpreting reported effect sizes within the context of the field. We recommend that NIH expect this information for key supporting data reported on animal research.

**RECOMMENDATION 4.2: NIH SHOULD EXPECT ALL VERTEBRATE AND CEPHALOPOD ANIMAL RESEARCH TO INCLUDE THE ARRIVE 2.0 ESSENTIAL 10 AT THE PUBLICATION STAGE.**

The ARRIVE guidelines 2.0 were published during this working group's activities [\[50, 51\]](#). In this update to the ARRIVE guidelines, authors improved the clarity of the original guidelines and prioritized them into two groups: the ARRIVE Essential 10 and the ARRIVE Recommended Set. The Essential 10 describes the information that the authors describe as "the basic minimum to include in a manuscript, as without this information, reviewers and readers cannot confidently assess the reliability of the findings presented." This list of information includes specific details about comparison groups and the experimental units, sample sizes, inclusion and exclusion criteria, randomization, blinding, outcome measures, statistical analysis methods, animals used, experimental procedures, descriptive statistics, and effect sizes and confidence intervals. We recommend that NIH expect that all manuscripts reporting on vertebrate and cephalopod research supported by NIH include the ARRIVE Essential 10.

**RECOMMENDATION 4.3: NIH SHOULD ENCOURAGE AND SUPPORT WORK TO BETTER UNDERSTAND, MONITOR, RECORD, AND REPORT IMPORTANT EXTRINSIC FACTORS RELATED TO ANIMAL CARE THAT MAY IMPACT RESEARCH RESULTS.**



**SUB-RECOMMENDATION 4.3A: NIH SHOULD PROVIDE EDUCATION ABOUT THE IMPORTANCE OF EXTRINSIC FACTORS TO THE RESEARCH COMMUNITY, PROVIDE A METHOD TO REPORT SUCH FACTORS, AND INCENTIVIZE PILOT STUDIES TO FURTHER IDENTIFY WHICH EXTRINSIC FACTORS ARE IMPACTFUL TO REPRODUCIBILITY.**

**SUB-RECOMMENDATION 4.3B: NIH SHOULD ESTABLISH A TASK FORCE TO IMPLEMENT THE CATALOGING OF EXTRINSIC FACTORS AS DATA FROM PILOT STUDIES ARE GATHERED.**

**SUB-RECOMMENDATION 4.3C: NIH SHOULD DEDICATE FUNDS FOR CONTROLLED RANDOMIZED TRIALS TO TEST THE EFFECT OF POTENTIALLY HIGH-VALUE EXTRINSIC FACTORS IDENTIFIED FROM PILOT STUDIES AND TASK FORCE RECOMMENDATIONS.**

Obtaining reproducible results in animal studies requires researchers to undertake a critical evaluation of all aspects of the study design that have an impact on experiences of the animals themselves. To reproduce animal experiments requires that researchers document and share critical information inherent to the animals (e.g., species, strain, sex) but also those extrinsic factors of the animals' environment (e.g., ambient temperature, microbiota, lighting levels) that systematically influence the experimental outcomes.

Much remains unknown about exactly which extrinsic factors in animal studies should be documented, with what temporal granularity documentation is needed, and how documentation should be shared for a study to be methodologically reproducible. Furthermore, no systematic characterization of the effects of various extrinsic conditions on different biological factors has been conducted. Without a better understanding of the impact of extrinsic factors on specific animal models or diseases, research outcomes will continue to suffer from unexplained differences both between and within laboratories and studies. We therefore recommend that NIH dedicate funds for the investigation of high-value extrinsic factors.

To raise awareness and understanding of the importance of extrinsic animal care factors, we recommend that NIH commit to educating investigators on appropriate animal data recording and reporting, which can occur in partnership with animal resource experts at their institutions. Investigators should be aware of how to obtain environmental and husbandry data points and to present impacts of deviations to standard housing and husbandry variables or standard operating procedures when necessary. Importantly, in order not to overburden investigators, much of the data regarding animal environmental conditions are currently tracked within existing animal program records (e.g., AAALAC program descriptions, daily housing room checklists) and can be provided to research teams by those involved in the delivery of animal care, including attending veterinarians or other animal care staff, Institutional Animal Care and Use Committee (IACUC) administrators, and grants and research program leadership. With consistent access to records of extrinsic factors, investigators can retain key information for their data files, and if deviations from expected outcomes occur, these can be explored, addressed, and reported in research findings per the ARRIVE 2.0 guidelines [\[50, 51\]](#).

One potential avenue for the disclosure of relevant extrinsic factors might be within the NIH RPPRs. NIH should consider how reporting key extrinsic factors in RPPRs, especially in a computable format, might aid in compiling information regarding the importance of these factors in experimental outcomes and/or facilitate sharing of these data more broadly. NIH should maintain a website or similar cloud-based resource to standardize the documentation of extrinsic factors and permit sharing of extrinsic factor data to the scientific community.

We also recommend that NIH develop a task force or working group to critically assess which extrinsic factors should be cataloged, identify how information could be stored and retained, and develop and fund the necessary technology for investigators to report these extrinsic factors so that they are computable and harvestable for data analysis. In addition, materials to educate and incentivize investigators about what factors should be reported and recorded, how to handle changes in conditions over the course of the award, and how and why to report this information in publications should be provided.

The task force should explicitly include end-users from the community of laboratory animal veterinary specialists, preclinical animal researchers who represent different animal systems, scientific publishers, and experts in machine learning and artificial intelligence to develop tools for assessing metadata. The charges of the task force should include the following:

- Determine how data are to be recorded, formatted, summarized, stored securely, and accessed/mined
- Identify benefits, costs, and possible burdens involved in recording extrinsic factors
- Assess what measures of impact can and should be systematically recorded
- Review and advise incentives and enforcement mechanisms to ensure accurate reporting

As a future goal, we recommend that NIH dedicate funding to study the potential impact of animal housing and husbandry environment on different conditions, such as behavior and the microbiome. As part of this project, NIH should:

- Establish goals for collecting data and how outcome variables are influenced by animal facility environmental factors (e.g., noise, vibrations, light, temperature)
- Specify or create funding programs that provide both a vehicle for data to be collected and approaches to analyzing data from the outcomes, with the resulting information published for public access

**RECOMMENDATION 4.4: NIH SHOULD PROVIDE SUPPORT FOR DOCUMENTING LARGER AND LONGER-LIVED ANIMALS' LONGITUDINAL EXPERIMENTAL, MEDICAL, AND HUSBANDRY HISTORIES.**

**SUB-RECOMMENDATION 4.4A: NIH SHOULD FORMALIZE FUNDING MECHANISMS TO LONGITUDINALLY RECORD AND MANAGE ANIMAL-LEVEL EXPERIMENTAL, MEDICAL, AND HUSBANDRY HISTORY METADATA FOR LARGER AND LONGER-LIVED ANIMALS.**

**SUB-RECOMMENDATION 4.4B: NIH SHOULD IDENTIFY MINIMAL ANIMAL-LEVEL EXPERIMENTAL, MEDICAL, AND HUSBANDRY HISTORY METADATA THAT WOULD BE LONGITUDINALLY RECORDED.**

**SUB-RECOMMENDATION 4.4C: NIH SHOULD ENCOURAGE THE SHARING OF ANIMAL-LEVEL EXPERIMENTAL, MEDICAL, AND HUSBANDRY HISTORY.**

With long-lived species, it is often the case that the group of people working with a particular animal will not remain constant across the animal's lifetime. It is therefore essential to encourage and support good record keeping for each animal. The community of researchers using larger and long-lived species must coalesce around the goal of establishing standards for maintaining useful records not only of the experimental histories of each animal but also of their medical histories and of the locations and characteristic of their housing. Medical records for larger long-lived species generally are already being kept by veterinarians and often by researchers as well. These records should remain with each animal

even if transferred. NIH can assist by bringing stakeholders together to identify minimal animal-level experimental, medical, and husbandry history metadata that would be longitudinally recorded. This additional record keeping improves methodological reproducibility and provides additional context when interpreting results. This documentation should be financially supported through NIH funding. Finally, NIH should communicate to researchers who use large animals the importance in sharing animal level experimental, medical, and husbandry history metadata. This documentation is in the hands of the researchers and veterinarians who work with the animals.

## Theme 5: Measure the Costs and Effectiveness of Efforts to Improve Rigor, Transparency, Reproducibility, and Translatability

The recommendations in this report have the potential to reshape the way we use animal models in biomedical research over the coming decade. Taken together, they compose a roadmap, with numerous recommendations structured for staged implementation ([Appendix 4](#)). We expect that the process of implementation will evolve and adjust in midcourse, informed by rigorous evaluation of both costs and effectiveness across different parts of the diverse NIH animal research portfolio.

While uncertainty is inherent in science, one certainty is that with few exceptions modern rigorous science requires financial support. As is true in many areas of life, scientists underestimate costs and time. A recent effort to test replicability of experiments reported in 50 high-impact science papers needed to be scaled back by more than a third because costs turned out to be much higher than expected [\[58\]](#). Likewise, most changes in the oversight and conduct of animal research outlined in the four preceding themes have cost implications, with increased financial costs to NIH being easy to imagine in concept yet difficult to predict in numbers. The changes recommended in this report will also bring opportunity costs. That is, given a finite NIH budget, increased animal costs needed to improve rigor will likely create opportunity costs for non-animal portions of the NIH enterprise. And within animal research, the financial costs of greater rigor applied in preclinical animal studies could lead to fewer grants or, alternatively, to grants with diminished scope.

We anticipate that counterbalancing these immediate increased costs will be longer-term savings. This will stem from outcomes such as increased scientific success rates and an improved efficiency resulting from building upon more rigorous findings. For example, if rigor is enhanced for research moving to the clinical research sphere, it is reasonable to expect there will be fewer failures of translation. These cost savings will open new scientific opportunities, and the savings from improved efficiency can be reinvested. As with costs, however, the magnitude of these savings is difficult to predict quantitatively.

NIH will therefore need to devise and implement ways to quantify costs, in dollars and in research opportunity, and to measure savings and success (e.g., the extent of improvement in rigor) for the changes it implements in pursuit of improved rigor. This perceived need prompted us to conduct preliminary integrative analyses of scientific rigor and grant economics by leveraging an existing set of more than 4,000 published research papers that had been coded for measures of rigor [\[59\]](#). These initial analyses suggested that enhanced rigor, especially sample size estimation, may indeed be associated with increased grant costs ([Appendix 5](#)). This pilot study also highlighted challenges with obtaining the kinds and amounts of data needed, including their computability, and drew attention to interpretive difficulties in discriminating between correlative and causal associations of rigor signatures with costs.

Our overall conclusion is that NIH should proactively design and lead evaluation of actions it takes in pursuit of rigor and use the results to hone its oversight practices and policies. Key activities will include identifying critical computable data and providing the means to obtain it, making specific plans for analysis, and appropriately evolving its ongoing actions based on interim results. While it is important for NIH to lead, we do not imagine that evaluation is a job that NIH can or should do alone. The research community should participate in the evaluative process from its design phase through the end analyses.

**RECOMMENDATION 5.1: NIH SHOULD DEVELOP AN EVALUATION PROGRAM TO ASSESS THE PROGRESS IN IMPLEMENTING THE REPORT RECOMMENDATIONS, THEIR EFFECTS ON NIH AND THE RESEARCH COMMUNITY, AND CHALLENGES THAT ARISE IN IMPLEMENTING RECOMMENDATIONS.**

Measuring improvements in quality is not a trivial activity. Decisions made by the NIH in support of the field of reproducibility research will have broad and far-reaching consequences. We were mindful of this and balanced our recommendations with a strong directive to NIH that implementation be staged, thoughtful, and periodically evaluated. As pilot studies recommended in this report or independently undertaken are completed and recommendations are implemented, NIH must be committed to ongoing review of the efforts. Detailed evaluation plans should be developed to ensure, as much as possible, that policies and strategies are having the intended effects. Considering that implementation of some recommendations may take significant time and resources, transparency and accountability are essential. It may be necessary for NIH to adjust course when implementing a recommendation outlined in this report. When such cases arise, NIH should use the accumulated evidence and work with stakeholders to modify strategies to achieve the desired goals.

**RECOMMENDATION 5.2: NIH SHOULD EXTERNALLY SUPPORT AND INTERNALLY CONDUCT ANALYSES ON ELEMENTS OF RIGOR AND TRANSPARENCY IN GRANT APPLICATIONS AND PUBLICATIONS TO EXAMINE THEIR FINANCIAL COSTS, OPPORTUNITY COSTS, AND IMPACT ON PORTFOLIO BALANCE.**

**SUB-RECOMMENDATION 5.2A: NIH SHOULD IDENTIFY AND COLLECT COMPUTATIONALLY EXTRACTABLE INFORMATION FROM GRANT PROPOSALS AND REPORTS ON POTENTIALLY IMPORTANT VARIABLES, INCLUDING PUBLICATION METRICS, METHODOLOGICAL RIGOR, FUNDING, INVESTIGATOR CAREER STAGE, INVOLVEMENT OF STATISTICIANS, EXPERIMENTAL DESIGN DESCRIPTIONS, AND NUMBERS AND SPECIES OF ANIMALS AND CONDUCT EXTENSIVE ANALYSES OF THESE DATA.**

**SUB-RECOMMENDATION 5.2B: NIH SHOULD ALLOW APPLICANTS TO INCLUDE TEXT IN THE BUDGET JUSTIFICATION SECTION ON HOW PROJECTED ANIMAL BUDGETS ARE LINKED TO EFFORTS TO ENHANCE TRANSPARENCY, RIGOR, AND REPRODUCIBILITY.**

NIH should support research on the resource implications of replication of animal research and of the implementation of measures to enhance rigor, transparency, and translatability. Through its evaluation of efforts aimed at improving rigor, NIH should build a foundation of evidence and analysis methods to enable ongoing monitoring and evolution of its rigor and reproducibility practices. Overall, NIH should monitor trends and changes in the nature, amount, and costs of its animal research model portfolio. Additional attention should be paid to changes in portfolio balance (e.g., between hypothesis generating and hypothesis testing work; between discovery, preclinical and clinical work.) In addition, NIH and/or others should be charged to specifically design and conduct analyses on sets of publications whose level of rigor has been or could be systematically assessed to extend and report how the level of rigor and other signatures of scientific quality are associated with other pertinent variables such as investigator career stage, involvement of statisticians, experimental design descriptions in grant proposals, and use

of registration. As a starting point, NIH should define what additional data on animal research it needs to collect and what high-priority questions it should ask to quantify financial costs and opportunity costs associated with higher levels of rigor. We discussed several entry points listed below. We note that a standing working group composed of all stakeholders with appropriate technical expertise could be chartered to aid NIH in designing and executing its evaluations:

- *Sample size:* More rigorous prospective sample size estimation will likely lead to increases in the number of required animals. Previous work has shown that scientists rarely report sample size estimation, and many if not most experiments may be substantially underpowered [60, 61]. If each experiment requires more animals, grant costs may increase, especially if the number of experiments per grant remains unchanged. There may need to be a culture shift at all levels in which each grant is expected to report fewer experiments.
- *Conduct of experiments:* Increasing rigor may require investments of more time and financial resources devoted to changes in the setup, conduct, and monitoring of experiments to include more detailed reporting and/or control of extrinsic factors (including feed, microbiome, housing, and climate), greater use of pre-application scientific review (see Recommendation 1.2), more intensive preparation and review of documents, increased use of multicenter protocols, and greater use of contract research organizations. NIH or its appointed study group should assess what data could/should be obtained to objectively measure these effects.
- *Model selection:* There may be a shift toward developing and using multiple animal systems, each designed to better answer a specific set of scientific questions or to model specific aspects of a human disease. Additionally, as researchers focus on selecting models that answer specific research questions, especially for translation to humans, we may see greater use of larger animals, including non-human primates. For example, the American Heart Association reviewed the different causes of heart failure in various animal systems and recommended a shift away from rodent models after it found that critical features of heart failure in patients did not align with the current rodent model [34]. NIH should gather data on the number, costs, and effectiveness of adding new animal models, retiring others, and employing non-animal models.
- *Infrastructure:* Development of infrastructure to support quality studies, prospective study design and analysis documentation (e.g., prospective registration), and enhanced data management, curation, storage, and sharing are expected to initially increase the cost of animal use. These costs should be evaluated relative to possible gains in efficiency that may balance the infrastructure investment over time.
- *Spillover effects:* There may be other effects outside of the context of specific grant applications and awards. Some scientists may choose to abandon certain areas of research because they and their institutions perceive new rigor requirements as overly restrictive, burdensome, or expensive. Institutions that are less well-resourced may make policy decisions to de-emphasize research involving animal models, leading to a distortion that narrows investigator diversity and institutionally concentrates which scientists are able to do animal research.
- *Other challenges:* At both the micro and macro levels, it is difficult to determine costs linked with specific experiments. Papers describing NIH-funded science often reference support from multiple sources, including more than one grant and more than one funding body. While publication and grant databases include a great deal of administrative data and metadata, other

types of valuable information may be hard to glean and may require either manual curation or sophisticated text-based machine-learning approaches.

As these recommendations are implemented across NIH, it will be important to allow and even encourage applicants to include text in the Budget Justification section to address how projected budgets are linked to efforts to enhance transparency, rigor, and translatability. Additional justification can include costs associated with enhanced transparency, rigor, and translatability. This could include justification of statistical personnel or services, the costs associated with replication, and transparent documentation and prospective registration. The explanations should focus on how each budget item is required to achieve enhanced transparency, rigor, and reproducibility.

**RECOMMENDATION 5.3: NIH SHOULD IDENTIFY SCIENTISTS WHO DEMONSTRATE THE HIGHEST LEVELS OF TRANSPARENCY AND RIGOR TO HELP DEFINE ENTERPRISE BEST PRACTICES.**

During our deliberations, we heard from several presenters on the science of rigor. As part of many presentations, we saw examples of effective practices they have adopted to improve rigor and enhance transparency in their laboratories ([Appendix 6](#)). NIH should identify scientists at all career stages and across scientific disciplines who consistently demonstrate the highest transparency and rigor. They should be recognized explicitly for their contributions to best practices, and their lessons and best practices should be promulgated by NIH.

## Conclusion

Successfully coupling strong scientific rigor and transparency with human curiosity and imagination is foundational to high-quality science. The frontiers of biomedical science are advancing rapidly, driven by breakthroughs that include facile gene editing, big data mining, artificial intelligence, and high-resolution imaging. Such progress is transforming animal research and presenting more opportunities for translation into improved human health outcomes. Yet the ability to interrogate increasingly complex animal biology comes with formidable challenges—both long-standing and new—to improve and more fully implement the best practices of the scientific method. This equally requires that researchers respect the ethical imperatives of studying animal subjects. We therefore strongly urge that NIH and the biomedical community coalesce around the shared mission to improve the rigor and transparency of research by partnering to test, implement, evaluate, and refine recommendations made here. The roadmap we outline aims to spark multiple actions, some of which can begin almost immediately. Overall, it is intended to guide a process that will play out over the coming decade to deliver lasting improvements. NIH must regularly communicate with the wider scientific community and the general public about the successes, obstacles, surprises, and evolution of these efforts. Our working group concludes its efforts with optimism that it is the right moment for NIH and the research community to embark on this joint endeavor with commitment, care, and necessary resources. We believe that all stakeholders will be rewarded with an unprecedentedly robust chain of new knowledge and improved health outcomes.

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## Appendices

### Appendix 1—Glossary

**Confirmatory Research:** Hypothesis-testing research to test the validity of an a priori hypothesis.

**Early-Stage Preclinical Research:** Animal research to understand the basis of human biology, disease, or disorders and develop interventions. This NIH usage is inclusive, and we note that the industry uses the term more narrowly to mean research focused on assessing the efficacy of candidate therapeutics.

**Exploratory Research:** Hypothesis-generating research to clarify the exact nature of the problem to be solved.

**Extrinsic Factors:** Factors that have a direct impact on the experience of the research animal during the course of experimental phases, such as housing, husbandry, handling, feed, water, bedding, enrichment, caging type, and light cycles.

**Generalizability:** Whether the results of a study apply in other contexts of populations that differ from the original one.

**Inferential Reproducibility:** When researchers draw similar conclusions or make knowledge claims of a similar strength from either an independent replication of a study or a reanalysis of the original study. This is part of the process by which a scientific field decides which research claims or effects are to be accepted as true.

**Late-Stage Preclinical Research:** Research using animals to find out whether a treatment is likely to be efficacious, often done immediately before testing in humans.

**Methods/Methodological Reproducibility:** When researchers provide enough detail about study procedures and data that the same procedures could, in theory or in actuality, be exactly repeated. Transparency is a key component of methods reproducibility.

**Prospective Registration:** Creation of a permanent record of a study design, an analytic plan, and a primary outcome before the data are collected. Prospective registration allows retrospective insurance against selective reporting and outcome switching. The registered research plan can be embargoed for a limited time to protect intellectual property, and when the registration is published, it allows identification and mitigation of publication bias. Researchers who prospectively register their studies demonstrate their intent to prioritize rigorous reporting. Prospectively registered studies do not undergo peer review at the time of registration.

**Publication Bias:** A form of bias in which the outcome of a study influences the decision to publish its results, resulting in the prioritization of positive results and large effects over null or negative results. Despite the availability of a range of journals and publishing outlets that welcome studies with null and negative results, publication bias is documented, indicating that researchers' behavior and incentive systems contribute to its occurrence.

**Reduction:** Appropriately designed and analyzed animal experiments that are robust and reproducible and truly add to the knowledge base.

**Refinement:** Advancing animal welfare by exploiting the latest *in vivo* technologies and by improving understanding of the impact of welfare on scientific outcomes.

**Registered Reports:** A journal article type in which the detailed study protocol is submitted for peer review before the data are collected. Upon successful review, the journal guarantees publication of the article regardless of the study findings. Like prospective registration, Registered Reports mitigate selective reporting and outcome switching. In addition, the protocol peer review can help improve experimental design, for example, by ensuring that sample size and inclusion/exclusion criteria are predetermined and that proper measures are taken to mitigate experimental and analysis bias. Researchers who publish with this review workflow demonstrate their intent to prioritize rigor, and they secure a path to publication regardless of findings, which mitigates publication bias.

**Replacement:** Accelerating the development and use of models and tools, based on the latest science and technologies, to address important scientific questions without the use of animals.

**Results Reproducibility:** When researchers conduct an independent new study with procedures as close to the original study as possible and obtain the same results. This definition is similar to others' definitions of "replicability."

**Scientific Rigor:** The strict application of the scientific method to ensure unbiased and well-controlled experimental design, methodology, analysis, interpretation, and reporting of results.

**Translation:** Applying results from preclinical research, usually via late-stage preclinical animal studies, to justify, design, and execute trials in humans.

**Transparency:** Accessibility of information.

## Appendix 2—Outcomes of High Rigor and Managing Expectations of Statistical Analysis

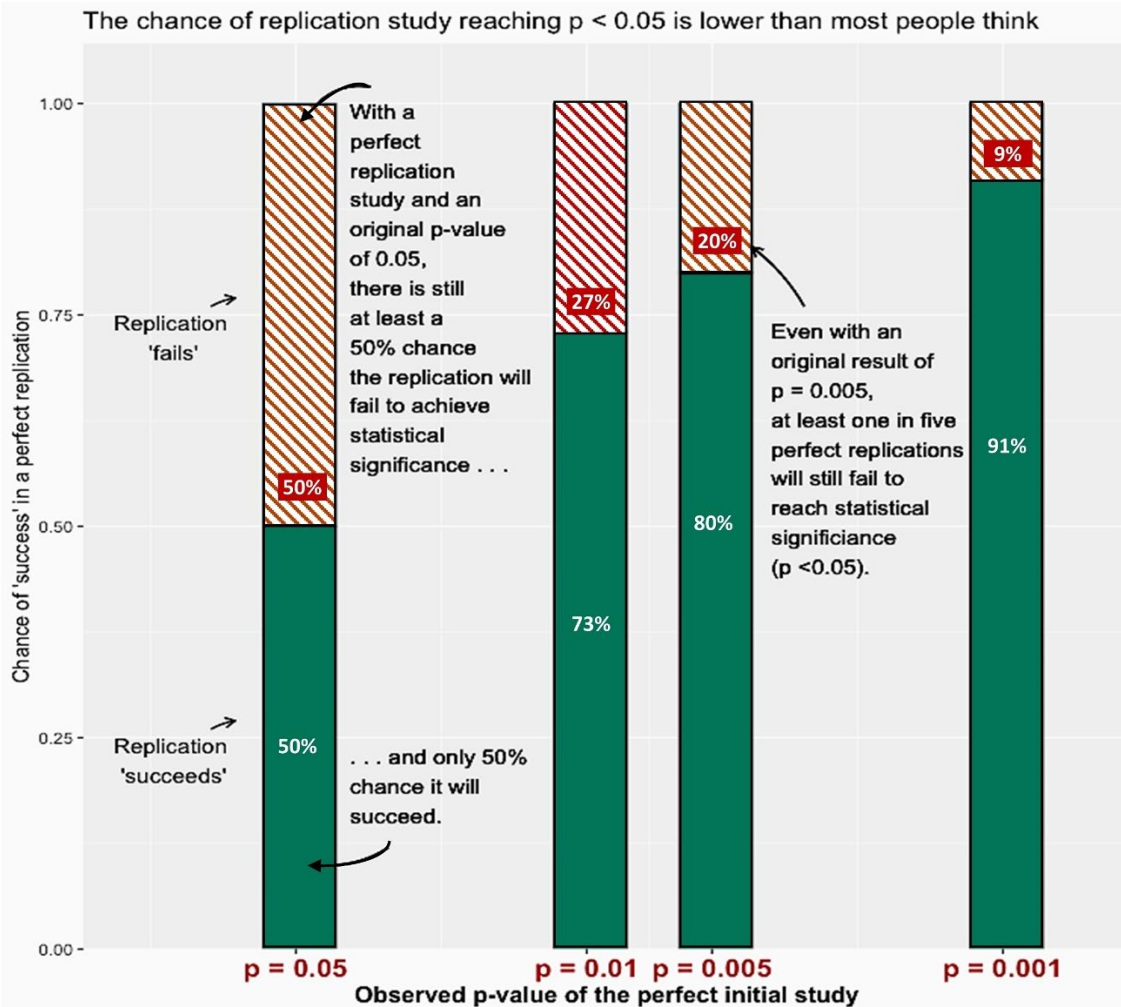


Figure 2. Managing Expectations of Statistical Analysis. Statistical significance is not enough to judge reproducibility. Given a statistically significant initial study, the chance of a replication “succeeding” (another statistical significance;  $p < 0.05$ ) is surprisingly low.

Much of good general statistical practice has been refined over the decades, particularly in clinical research [62-65]. Yet preclinical animal research has unique statistical needs and challenges, many of which have not been fully addressed by the statistical and animal research communities. For example, due to ethical considerations and resource constraints, animal researchers are under great pressure to minimize the number of animals in their studies. Yet studies with smaller sample sizes generally have lower statistical power, which leads to more false negatives (overlooking true effects) and more false positives (incorrectly declaring an effect to be statistically significant). To achieve adequate statistical power with the minimum number of animals, which is essential for confirmatory and late-stage preclinical experiments, researchers need to use appropriate experimental design and sample size calculations, but many animal researchers report lacking the knowledge or impetus to do so [66, 67].

Furthermore, although researchers may have more genetic and environmental control over animals than over humans in clinical research, they must still deal with the animal-to-animal variations that occur naturally and from induced injury and disease; this makes sample size calculations crucial for ensuring that experimental effects can be estimated with enough precision to be informative [68, 69]. Similarly, the preferential use of male animals has been a long-standing issue in animal experiments [70-72], which has led to a disadvantage for women by skewing our understanding of diseases and the development of potential therapies to disproportionately favor males. NIH instituted sex as a biological variable (SABV) in 2016, requiring inclusion of male and female subjects, which increased overall animal use and sample sizes intentionally to aid in translation and applicability to human disease; researchers and their funders must balance the importance of considering sex as a biological variable with its potential impact on statistical power and the ethical use of both sexes of animals in modeling human disorders [73].

Sources of confounding can also arise at virtually every stage of an animal experiment [74-76], such as room temperature [74-76], group housing [77, 78], lighting [74], cage environmental stability [79], and even the sex of the experimenter [80]. Preclinical animal research also lacks established analytic tools, such as the “minimally important difference” common in clinical trials or meta-analytical techniques that can handle the diversity of species and variations in animal study designs [81, 82].

Using animals to model human physiology and disease comes with well-known limitations. Yet even the best animal models are ineffective if the studies employing them also have improper experimental design, inappropriate data analysis, inadequate results reporting, or other statistical problems [36, 83-87]. One analysis, for example, found that 91% of reviewed studies in a particular field failed to use the correct experimental unit in their analyses, leading to incorrect claims that would be impossible to replicate [88]. Another analysis estimated that problems in study design, data analysis, and statistical reporting account for more than 50% of study errors that lead to irreproducibility in U.S. research [84].

## Appendix 3—Additional Context for Recommendation 1.3

### Proposed information to guide applicants in addressing each critical study design element

The following items are critical to ensure rigorous research by minimizing risk of bias and fostering transparency. Describe how these items will be addressed for each proposed study design in the application, alongside unambiguous descriptions of animals used, where applicable. These items are also expected to be reported appropriately when communicating study results.

- 1. Inclusion/exclusion criteria:** *Describe the criteria that will be used for inclusion or exclusion of samples or animals during the experiments and for data used in analysis.*
- 2. Sample size estimation:** *Provide planned sample sizes for each group and how they were derived.*
- 3. Data analysis plan:** *Describe plans for data analysis, including statistical methods as appropriate, designed to answer the proposed scientific questions.*
- 4. Blinding:** *Describe measures planned to blind the investigators during group allocation, the conduct of the experiment, and the analysis, where applicable. If none have been taken and blinding is not appropriate to the study design, provide justification.*
- 5. Randomization:** *Describe methods planned for random allocation to comparison groups and strategies for random sample processing and collection of data where applicable. Provide a rationale if a randomization scheme is not used.*

### Resources to support successful completion of additional page

A landing page of FAQs regarding rigor and reproducibility [currently exists at nih.gov](#), and the above resources could be added to this site. However, it is the recommendation of the subcommittee that this page be updated and linked to additional resources for preparation of grant applications.

To support applicants and facilitate satisfactory completion of details required in the additional page, we recommend that resources additionally describing these items be made readily available to applicants on the NIH Grants & Funding portal. Such resources could include:

- **NIH Rigor and Reproducibility Resources**

<https://www.nih.gov/research-training/rigor-reproducibility/resources>

- **NIH Principles and Guidelines for Reporting Preclinical Research**

<https://www.nih.gov/research-training/rigor-reproducibility/principles-guidelines-reporting-preclinical-research>

- **The Experimental Design Assistant (EDA)**

<https://journals.plos.org/plosbiology/article?id=10.1371/journal.pbio.2003779>

<https://www.nc3rs.org.uk/experimental-design-assistant-eda>

- **Nature Journals Reporting Summary**



<https://www.nature.com/documents/nr-reporting-summary-flat.pdf> [PDF-5.19M] As part of *Nature Research* reporting requirements: <https://www.nature.com/nature-research/editorial-policies/reporting-standards#reporting-requirements>

- **The ARRIVE Guidelines 2.0**

du Sert NP, Hurst V, Ahluwalia A, Alam S, Avey MT, Baker M, et al. The ARRIVE guidelines 2.0: updated guidelines for reporting animal research. *PLOS Biol*, 2020;18(7):e3000410. <https://doi.org/10.1371/journal.pbio.3000410>. Accessed June 25, 2021.

du Sert NP, Ahluwalia A, Alam S, Avey MT, Baker M, Browne WJ, et al. Reporting animal research: explanation and elaboration for the ARRIVE guidelines 2.0. *PLOS Biol*, 2020;18(7):e3000411. <https://doi.org/10.1371/journal.pbio.3000411>.

- **The PREPARE Guidelines**

Smith AJ, Clutton RE, Lilley E, Hansen KEA, Brattelid T. PREPARE: guidelines for planning animal research and testing. *Lab Anim*, 2018;52(2):135–141. <http://doi.org/10.1177/0023677217724823>. Epub 2017. PMID: 28771074; PMCID: PMC5862319.

- **Additional Reporting Guidelines**

Landis S, Amara S, Asadullah K, et al. A call for transparent reporting to optimize the predictive value of preclinical research. *Nature*, 2012;490:187–191. <https://doi.org/10.1038/nature11556>.

Chambers K, Collings A, Graf C, Kiermer V, Mellor DT, Macleod M, et al. Towards Minimum Reporting Standards for Life Scientists. *MetaArXiv*, April 30, 2019. <http://doi.org/10.31222/osf.io/9sm4x> (including reporting information framework [<https://osf.io/xfpn4/>] and elaboration document [<https://osf.io/xzy4s/>]).

## Appendix 4—Report Recommendation Implementation Gantt

As part of our recommendation planning, we developed a possible timeline for the NIH to consider when implementing our recommendations.

Actions	Q2 2021	Q3 2021	Q4 2021	Q1 2022	Q2 2022	Q3 2022	Q4 2022	Q1 2023	Q2 2023	Q3 2023	Q4 2023	Q1 2024	Q2 2024	Q3 2024	Q4 2024	Q1 2025	Q2 2025	Q3 2025	Q4 2025	Q1 2026	Q2 2026	Q3 2026	Q4 2026	Q1 2027	Q2 2027	Q3 2027	Q4 2027	Q1 2028	Q2 2028	Q3 2028	Q4 2028
<b>Theme 1: Improve Study Design and Data Analysis</b>																															
<b>Recommendation 1.1</b>																															
Sub-recommendation 1.1a	Development																														
Sub-recommendation 1.1b	Development																														
Sub-recommendation 1.1c	Funding																														
<b>Recommendation 1.2</b>	5-year development project																														
Sub-recommendation 1.2a	Planning (Phase 1 - current systems)																														
Sub-recommendation 1.2b	Phased Implementation (phase 1 - current systems)																														
Sub-recommendation 1.2c	Plan (phase 2 - new systems)																														
Sub-recommendation 1.2d	Phased Implementation (phase 2 - new systems)																														
Sub-recommendation 1.4a	Planning/Budgeting																														
Sub-recommendation 1.4b	Pilot FOA																														
Sub-recommendation 1.4c	FOAs																														
Sub-recommendation 1.4d	FOAs																														
<b>Recommendation 1.3</b>	Plan																														
<b>Recommendation 1.4</b>	Announce																														
	Evaluate																														
	Recs																														
<b>Theme 2: Address Incomplete Reporting and Questionable Research Practices</b>																															
<b>Recommendation 2.1</b>	Report																														
<b>Recommendation 2.2</b>	Planning																														
Sub-recommendation 2.2a	Phased Campaign - Education, Awareness																														
Sub-recommendation 2.2b	Phased Campaign - Education, Awareness & outcomes																														
	multiyear projects and evaluation																														
	Phase 2 Implementation																														
	multiyear projects and evaluation																														
	Phase 2 Implementation																														
	multiyear projects and evaluation																														
	Phase 2 Implementation																														
<b>Theme 3: Improve Selection, Design, and Relevance of Animal Models</b>																															
<b>Recommendation 3.1</b>	Planning																														
<b>Recommendation 3.2</b>	Development																														
<b>Recommendation 3.3</b>	Announce																														
<b>Recommendation 3.4</b>	Phased Implementation																														
Sub-recommendation 3.4a	Planning/Budgeting																														
Sub-recommendation 3.4b	Pilot FOA																														
<b>Recommendation 3.5</b>	Multiyear Projects; Evaluation; Planning and Budgeting for Phase 2																														
<b>Recommendation 3.6</b>	FOAs																														
	Multiyear Projects																														
	Plan																														
	Announce																														
	Phased Implementation																														
	Plan																														
	Announce																														
	Phased Implementation																														
	Report																														
	Phased Campaign - Education, Awareness																														
	Plan																														
	Announce																														
	Deliberations																														
	Recs																														
<b>Theme 4: Improve Methodological Documentation and Results Reporting</b>																															
<b>Recommendation 4.1</b>	Plan																														
<b>Recommendation 4.2</b>	Announce																														
<b>Recommendation 4.3</b>	Phased Implementation																														
Sub-recommendation 4.3a	Report																														
Sub-recommendation 4.3b	Plan																														
Sub-recommendation 4.3c	Announce																														
<b>Recommendation 4.4</b>	Phased Campaign - Education, Awareness																														
Sub-recommendation 4.4a	Plan																														
Sub-recommendation 4.4b	Announce																														
Sub-recommendation 4.4c	Deliberations																														
	Recs																														
	Planning/Budgeting																														
	FOAs; Planning/Budgeting for Phase 2																														
	FOAs; Multiyear Projects; Evaluation																														
	Multiyear Projects; Evaluation																														
	Multiyear Projects; Evaluation																														
	Multiyear Projects; Evaluation																														
	Multiyear Projects; Evaluation																														
	Planning																														
	Development																														
	Phased Implementation																														
	Plan and Deliberations																														
	Announce																														
	Phased Implementation																														
	Report																														
	Announce																														
<b>Theme 5: Measure the Costs and Effectiveness of Efforts to Improve Rigor, Transparency, Reproducibility, and Translatability</b>																															
<b>Recommendation 5.1</b>	Planning																														
<b>Recommendation 5.2</b>	Development																														
Sub-recommendation 5.2a	Phased Evaluation																														
Sub-recommendation 5.2b	Planning																														
<b>Recommendation 5.3</b>	Development																														
	Announcement																														
	Phased Implementation																														
	Internal and External Analyses																														
	Analysis and Development of Best Practices																														

## Appendix 5—Preliminary Appendix Report: Potential Financial Ramifications of Working Group Recommendations

Michael Lauer, Robyn Lee-Stubbs, Sarah Nusser, Regina Nuzzo, Eric Prager\*

\*Authors are listed in alphabetical order

May 23, 2021

### Introduction

While uncertainty is inherent in science, one certainty is that with few exceptions, modern rigorous science requires financial support. As is true in many areas of life, scientists underestimate costs and time. A [recent effort](#) to replicate the experiments reported in 50 high-impact science papers was scaled back because costs turned out to be much higher than expected.

The subcommittee engaged in three sets of activities:

- Developed a set of possible financial implications of substantive changes to the way that NIH funds and oversees research involving animal models.
- Conducted two preliminary analyses of scientific rigor and grant economics by leveraging data from publications in [Circulation Research](#) and [Stroke](#) of more than 5,000 published research papers that had been coded for measures of rigor. This preliminary report is based on the latter publication, because it included a larger sample with publications occurring over a wider range of years.
- Developed recommendations.

### Possible Financial Implications

Changes in the funding, oversight, and conduct of research involving animal models may lead to increased costs to the NIH (and therefore opportunity costs for the entire NIH-funded enterprise) for a number of reasons:

- *Numbers of animals:* Rigorous prospective sample size estimation will likely lead to substantial increases in the number of required animals. [Previous work](#) has shown that scientists rarely report sample size estimation and that many if not most [experiments may be substantially underpowered](#). If each experiment requires more animals, grant costs may increase, especially if the number of experiments per grant remains unchanged. In other words, there may need to be a shift in expectations whereby each grant (as well as each paper) reports fewer experiments, but each experiment will be properly powered.
- *Type of animals:* There may be a shift toward greater use of larger animals (including non-human primates) and/or a greater diversity of species used for each scientific question. For example, the [American Heart Association](#) described existing animal models for different causes of heart failure and recommended that there be a shift away from rodent models.

- *Conduct of experiments:* Changes in the setup, conduct, and monitoring of experiments include greater levels of control of living conditions (including feed, microbiome, housing, and climate), greater use of pre-IACUC scientific review, more intensive preparation and review of documents, increased use of multicenter protocols (such as the [Intervention Testings Program](#) of the National Institute on Aging), greater use of contract research organizations, and development of infrastructures needed to support replication studies, preregistration, and enhanced data management, curation, storage, and sharing.
- *Spillover effects:* There may be other effects that occur outside of the context of specific grant applications and awards. Some scientists may choose to abandon certain areas of research because they and their institutions perceive new rigor requirements as overly restrictive, burdensome, or expensive. Institutions that are less well-resourced may make policy decisions to de-emphasize research involving animal models, leading to an increased concentration of research resources, a trend that [some see as concerning](#). On the other hand, it can be argued that poorly designed (including underpowered) research is inherently [wasteful and unethical](#). Furthermore, greater rigor in preclinical research should lead to fewer efforts in the clinical research sphere to study the effects of candidate agents or devices that should never have been studied in the first place.
- *Other challenges:* At both the micro and macro levels, it is difficult to determine costs linked with specific experiments. Many papers describing NIH-funded science mention support from more than one source—including more than one grant and more than one funding body (e.g., NIH, nonprofit foundations, industry, academic institutions). Although publication and grant databases include a great deal of administrative data and metadata, other types of valuable information may be hard to glean, requiring either manual curation or sophisticated text-based machine-learning approaches.

#### Preliminary Analyses

F. Daniel Ramirez, a member of the ACD Working Group, kindly provided us with data based on an [analysis of rigor](#) of 4,162 cardiovascular articles published over 18 years. The articles were selected from three journals (*Stroke*, *Nature Medicine*, and *Science Translational Medicine*) that had implemented steps in 2011 to enhance rigor and from two “control” journals. All papers described experiments involving nonhuman mammals. Dr. Ramirez provided us with data for each paper, including whether there was reporting of randomization, blinding, sample size estimation, and sex used.

The data we received did not include paper ID numbers (e.g., Scopus, PubMed). We were able to match 4,001 papers (96%) with PubMed ID numbers. We used the [NIH SPIRES database](#) to link 2,348 papers to 6,073 unique NIH grants. We captured the complete history of each grant, including total years of funding up to the year of publication, total grant dollars, and total dollars during the most recent year of funding. All funding values were corrected for inflation using the [Biomedical Research and Development Price Index \(BRDPI\)](#) (2019 reference value).

In our effort to gain insights into possible fiscal ramifications of rigor recommendations that the ACD WG might make, we leverage three types of data:

- Article metrics: [relative citation ratio](#) (RCR), and the [approximate potential to translate](#) (APT), namely the likelihood that a paper will be cited in a guideline or clinical trial publication
- Methodological rigor as coded by [Ramirez et al.](#): randomization, blinding, and sample size estimation, as well as reporting of sex

- Grant measures: number of grants cited, grant years of prior support, total grant funding up to publication (in inflation-adjusted dollars), most recent grant funding, and use of certain specific mechanisms (U, G, M, P, and training)

We use these data to consider three possible desirable outcomes:

- For research conducted and reported with methodological rigor, we use data from [Ramirez et al.](#)
- For research that is translatable, at least in some instances, we use APT data from the [NIH Office of Portfolio Analysis](#).
- For research that is influential, though this may be particularly controversial, we use RCR data from the [NIH Office of Portfolio Analysis](#).

NIH Funding and Measures of Article Rigor, Influence, and Translatability

We were able to find PubMed ID numbers for 2,348 NIH-funded papers and 1,653 papers funded entirely from other sources.

The NIH Office of Portfolio Analysis has developed a tool by which one can map a set of papers according to the “Triangle of Biomedicine” of [Weber](#). The triangle uses [Medical Subject Heading](#) terms to classify papers according to axes of molecular/cellular, animal, or human focus.

Figures 3 and 4 show the triangles for NIH- and non-NIH-funded papers. As expected, all papers gravitate toward the animal corner. NIH- and non-NIH-funded papers appear to be similar by this scheme.

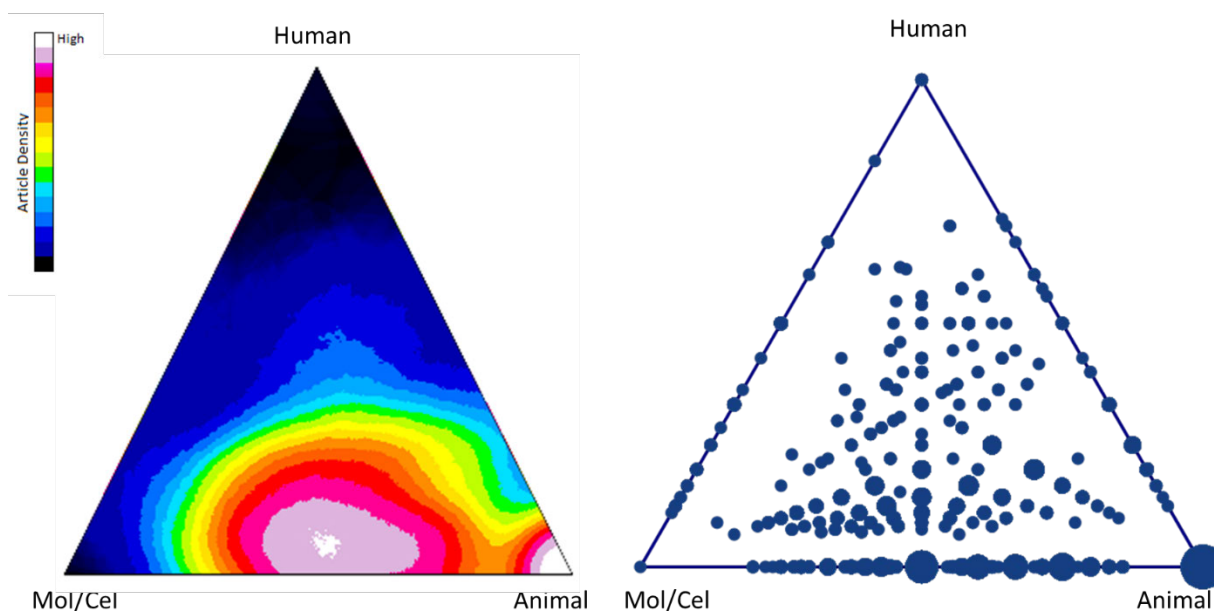


Figure 3: Triangle of Biomedicine for 2,348 NIH-funded Papers from Ramirez et al. Heat map or dot plot maps of science to visually depict the structure of literature by showing the relationships among different fields of science. The corners of the triangle correspond to animal (A), cellular or molecular (C), and human (H) research. The heat map or size of the circle is proportional to the number of articles in that discipline.

[Table 1](#) shows article, influence, translatability, and rigor measures according to source of funding. We considered papers to be of higher influence if they had an RCR greater than 3 and to be of higher

translatability if the approximate potential to translate was greater than 50%. We define “rigor points” as the number of rigor measures (randomization, blinding, and power) described and define a paper as “more rigorous” if at least one of these rigor measures is present. We considered articles to be “most rigorous” if all three measures of rigor—randomization, blinding, and sample size estimation—were present. NIH-funded papers were more likely to be published in general journals (e.g., *Nature Medicine*, *Science Translational Medicine*) and use mouse models and were less likely to use rat models. Influence measures were higher for NIH-funded papers, while translatability measures were similar. NIH-funded papers were less likely to report randomization and blinding but were more likely to report sample size estimation. Even so, only 11% of NIH-funded paper reported formal sample size estimation.

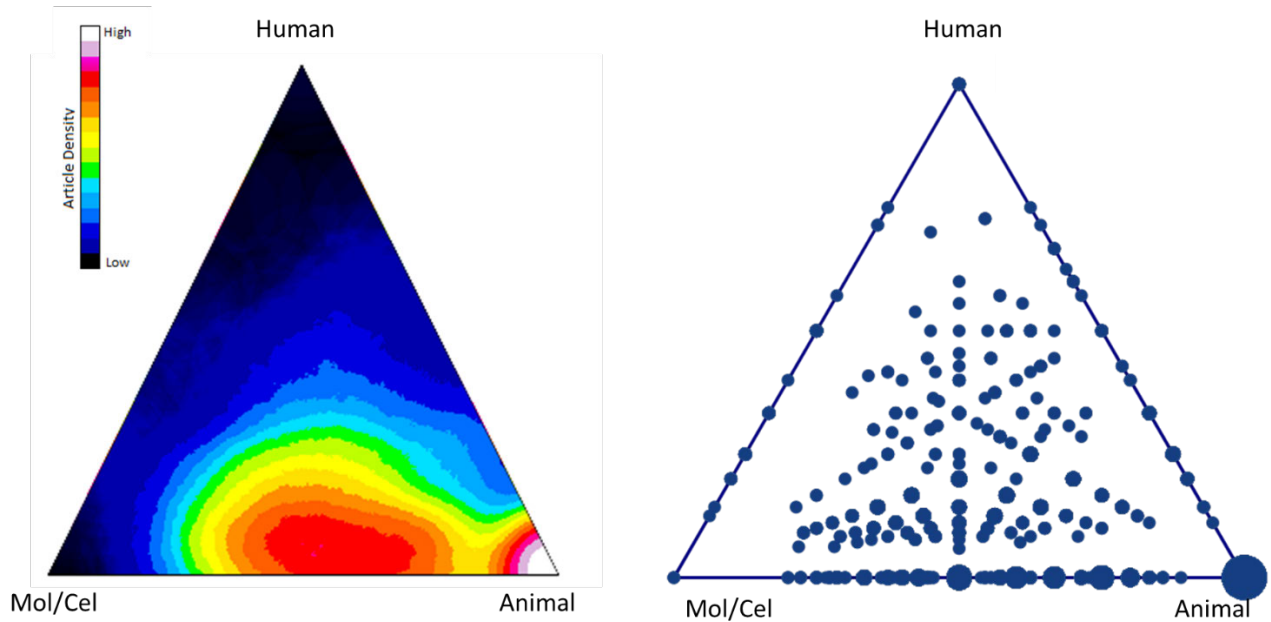


Figure 4: Triangle of Biomedicine for 1,653 Non-NIH-funded Papers from Ramirez et al. Heat map or dot plot maps of science to visually depict the structure of literature by showing the relationships among different fields of science. The corners of the triangle correspond to animal (A), cellular or molecular (C), and human (H) research. The heat map or size of the circle is proportional to the number of articles in that discipline.



Table 1: Characteristics of Articles According to Source of Funding

Characteristic/Funding	NIH-Funded		Not NIH-Funded
Total N (%)	2,348 (58.7)		1,653 (41.3)
Model	Mouse	1,680 (71.6)	1,003 (60.7)
	Rat	328 (14.0)	393 (23.8)
	Combination	127 (5.4)	119 (7.2)
	Primate	52 (2.2)	29 (1.8)
	Rabbit	53 (2.3)	26 (1.6)
	Dog	42 (1.8)	30 (1.8)
	Pig	42 (1.8)	27 (1.6)
	Other	24 (1.0)	26 (1.6)
Journal	<i>Stroke</i>	465 (19.8)	518 (31.3)
	<i>Nat Med</i>	562 (23.9)	326 (19.7)
	<i>Sci Transl Med</i>	520 (22.1)	281 (17.0)
	<i>Circulation</i>	422 (18.0)	322 (19.5)
	<i>Circ Res</i>	379 (16.1)	206 (12.5)
Intervention Journal	Yes	1,547 (65.9)	1125 (68.1)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	1,082 (46.1)	607 (36.7)
Relative Citation Ratio	Median (IQR)	2.9 (1.8 to 5.4)	2.5 (1.5 to 4.5)
Higher Influence	Yes	1142 (48.6)	682 (41.3)
Approximate Potential to Translate	Median (IQR)	0.5 (0.2 to 0.8)	0.5 (0.2 to 0.8)
Higher Translatability	Yes	1126 (48.0)	769 (46.5)
Randomization	Yes	729 (31.0)	581 (35.1)
Blinding	Yes	935 (39.8)	725 (43.9)
Sample Size Estimation	Yes	258 (11.0)	114 (6.9)
Rigor Points	0	1,080 (46.0)	701 (42.4)
	1	744 (31.7)	546 (33.0)
	2	394 (16.8)	344 (20.8)
	3	130 (5.5)	62 (3.8)
More Rigorous	Yes	1,268 (54.0)	952 (57.6)
Most Rigorous	Yes	130 (5.5)	62 (3.8)

Article Metrics, Rigor Measures, and Grant Characteristics Among NIH-funded Papers

[Table 2](#) shows article metrics, rigor, and grant measures for the 4,001 papers according to the animal model. As might be expected, papers reporting on primate models and combination models were associated with greater grant costs.

In [Tables 3, 4, and 5](#), we break down article metrics and grant measures according to the specific type of rigor measure. Of note, sample size estimation was associated with greater grant costs ([Table 5](#)). [Figure 5](#) shows the distribution of the most recent grant dollars according to whether there was sample size estimation. Power calculation was the least common measure of rigor, though it appeared to be associated with greater costs. The box plots show, as might be expected given the sample imbalance, a wider distribution and a higher mean for papers without a power calculation.

Table 2: Characteristics of Articles and Associated Grants According to the Animal Model

Characteristic/Model		Mouse	Rat	Combination	Primate	Rabbit	Dog	Pig	Other
Total N (%)		1,680 (71.6)	328 (14.0)	127 (5.4)	52 (2.2)	53 (2.3)	42 (1.8)	42 (1.8)	24 (1.0)
Relative Citation Ratio	Median (IQR)	3.1 (1.8 to 5.7)	2.4 (1.6 to 3.8)	3.8 (2.3 to 6.2)	3.0 (2.0 to 5.8)	2.1 (1.2 to 3.1)	2.0 (1.5 to 3.6)	2.6 (1.7 to 4.6)	2.4 (1.7 to 4.3)
Higher Influence	Yes	863 (51.4)	117 (35.7)	79 (62.2)	26 (50.0)	15 (28.3)	14 (33.3)	18 (42.9)	10 (41.7)
Approximate Potential to Translate	Median (IQR)	0.5 (0.2 to 0.8)	0.5 (0.2 to 0.8)	0.8 (0.5 to 0.8)	0.8 (0.5 to 0.8)	0.5 (0.2 to 0.8)	0.5 (0.5 to 0.8)	0.5 (0.2 to 0.8)	0.5 (0.2 to 0.8)
Higher Translatability	Yes	816 (48.6)	131 (39.9)	73 (57.5)	33 (63.5)	25 (47.2)	20 (47.6)	19 (45.2)	9 (37.5)
Intervention Journal	Yes	1060 (63.1)	260 (79.3)	95 (74.8)	50 (96.2)	35 (66.0)	10 (23.8)	23 (54.8)	14 (58.3)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	874 (52.0)	36 (11.0)	90 (70.9)	47 (90.4)	8 (15.1)	3 (7.1)	17 (40.5)	7 (29.2)
Randomization	Yes	471 (28.0)	122 (37.2)	58 (45.7)	14 (26.9)	19 (35.8)	13 (31.0)	21 (50.0)	11 (45.8)
Blinding	Yes	660 (39.3)	156 (47.6)	51 (40.2)	7 (13.5)	21 (39.6)	9 (21.4)	20 (47.6)	11 (45.8)
Sample Size Estimation	Yes	192 (11.4)	19 (5.8)	24 (18.9)	5 (9.6)	6 (11.3)	2 (4.8)	8 (19.0)	2 (8.3)
Rigor Points	0	791 (47.1)	132 (40.2)	47 (37.0)	31 (59.6)	26 (49.1)	28 (66.7)	14 (33.3)	11 (45.8)
	1	548 (32.6)	107 (32.6)	39 (30.7)	16 (30.8)	13 (24.5)	5 (11.9)	13 (31.0)	3 (12.5)
	2	248 (14.8)	77 (23.5)	29 (22.8)	5 (9.6)	9 (17.0)	8 (19.0)	9 (21.4)	9 (37.5)
	3	93 (5.5)	12 (3.7)	12 (9.4)	0 (0.0)	5 (9.4)	1 (2.4)	6 (14.3)	1 (4.2)
More Rigorous	Yes	889 (52.9)	196 (59.8)	80 (63.0)	21 (40.4)	27 (50.9)	14 (33.3)	28 (66.7)	13 (54.2)
Number of Grants	Median (IQR)	4.0 (2.0 to 6.0)	3.0 (2.0 to 3.0)	4.0 (2.0 to 6.0)	4.0 (2.0 to 6.0)	2.0 (2.0 to 4.0)	2.5 (2.0 to 4.0)	3.5 (2.0 to 5.0)	2.0 (2.0 to 4.0)
Grant-Years of Funding	Median (IQR)	28.0 (13.0 to 50.0)	16.0 (7.0 to 28.0)	26.0 (10.5 to 51.0)	23.0 (9.8 to 48.0)	20.0 (8.0 to 25.0)	12.5 (9.0 to 26.0)	19.0 (11.5 to 37.8)	12.0 (5.0 to 27.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	15.5 (4.4 to 48.0)	4.1 (1.8 to 12.0)	18.6 (5.4 to 40.7)	56.5 (9.2 to 150.4)	7.2 (2.8 to 15.2)	11.0 (1.9 to 17.8)	9.0 (3.8 to 22.6)	4.4 (1.4 to 7.2)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	1.9 (0.6 to 5.4)	0.6 (0.3 to 1.2)	2.3 (0.6 to 5.4)	8.1 (1.2 to 16.8)	0.8 (0.4 to 2.6)	0.8 (0.3 to 3.4)	1.1 (0.4 to 2.6)	0.5 (0.3 to 0.9)
P, U, M, or G Mechanism	Yes	975 (58.0)	140 (42.7)	76 (59.8)	42 (80.8)	32 (60.4)	20 (47.6)	20 (47.6)	4 (16.7)
Training Grant	Yes	381 (22.7)	21 (6.4)	28 (22.0)	10 (19.2)	4 (7.5)	5 (11.9)	5 (11.9)	1 (4.2)



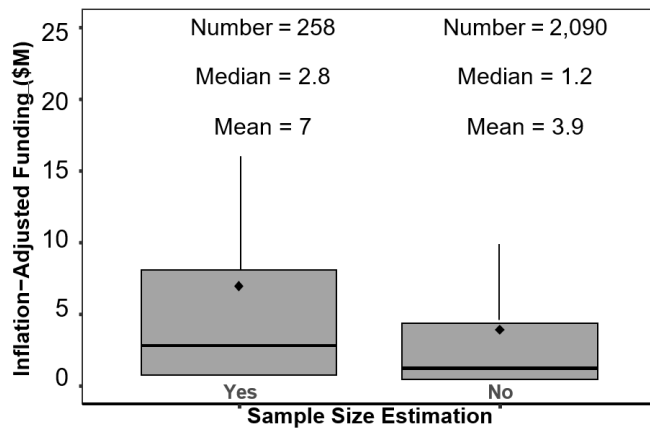


Figure 5: Distribution of Recent Grant Funding According to Whether There Was Sample Size Estimation

In [Table 6](#), we compare paper and grant history measures according to whether the paper appears to have a higher translatability, which we define as a value greater than 50%. Articles with higher translatability had higher relative citation ratios, higher rates of randomization, higher rates of sample size estimation, and higher grant costs. In [Table 7](#), we compare article and grant history measures according to whether the paper appears to have a higher level of influence, which we define as an RCR greater than 3. Higher influence papers had higher rates of sample size estimation and higher grant costs.

#### Article and Grant Measures According to Sex

[Table 8](#) shows article metrics, rigor, and grant measures according to the sex of animals described in the papers. Papers that reported on experiments involving males and females had higher grant costs.

#### Exploratory Regression Analyses

[Table 9](#) shows exploratory [logistic regression analyses](#) of four paper-based outcomes, namely randomization, blinding, sample size estimation, and reporting of any of these three types of rigor (what we have called “more rigorous”). None of the models explain the outcomes well. There may be a weak association between grant funding and higher odds of sample size estimation.

At this point, we should keep in mind some important limitations:

- The data are limited to cardiovascular papers published between 2000 and 2017.
- Few papers reported on sample size estimation.
- Other measures of interest likely require manual curation.
- Grant acknowledgments are not standardized.
- While sample size calculation appears to be associated with increased costs (as might be expected), grant costs explain little of the variance.
- These analyses should be considered only preliminary; they present an example of what might be possible but should not be construed as suggesting or reaching any conclusions.

Table 3: Characteristics of Articles and Associated Grants According to Whether There Was Randomization

Characteristic		Randomization	No Randomization
Total N (%)		729 (31.0)	1,619 (69.0)
Relative Citation Ratio	Median (IQR)	2.9 (1.8 to 5.5)	2.9 (1.8 to 5.3)
Higher Influence	Yes	356 (48.8)	786 (48.5)
Approximate Potential to Translate	Median (IQR)	0.8 (0.2 to 0.8)	0.5 (0.2 to 0.8)
Higher Translatability	Yes	378 (51.9)	748 (46.2)
Intervention Journal	Yes	615 (84.4)	932 (57.6)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	416 (57.1)	666 (41.1)
Number of Grants	Median (IQR)	3.0 (2.0 to 5.0)	3.0 (2.0 to 6.0)
Grant-Years of Funding	Median (IQR)	25.0 (12.0 to 46.0)	25.0 (11.0 to 44.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	13.1 (4.1 to 53.1)	11.3 (3.3 to 34.1)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	1.5 (0.5 to 5.7)	1.3 (0.4 to 4.3)
P, U, M, or G Mechanism	Yes	427 (58.6)	882 (54.5)
Training Grant	Yes	140 (19.2)	315 (19.5)

Table 4: Characteristics of Articles and Associated Grants According to Whether There Was Blinding

Characteristic		Blinding	No Blinding
Total N (%)		935 (39.8)	1,413 (60.2)
Relative Citation Ratio	Median (IQR)	2.8 (1.8 to 5.2)	3.0 (1.8 to 5.5)
Higher Influence	Yes	438 (46.8)	704 (49.8)
Approximate Potential to Translate	Median (IQR)	0.5 (0.2 to 0.8)	0.5 (0.2 to 0.8)
Higher Translatability	Yes	443 (47.4)	683 (48.3)
Intervention Journal	Yes	687 (73.5)	860 (60.9)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	425 (45.5)	657 (46.5)
Number of Grants	Median (IQR)	3.0 (2.0 to 6.0)	3.0 (2.0 to 6.0)
Grant-Years of Funding	Median (IQR)	24.0 (10.0 to 44.0)	25.0 (11.0 to 46.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	10.8 (3.3 to 36.7)	12.5 (3.7 to 41.1)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	1.1 (0.4 to 4.2)	1.6 (0.5 to 5.1)
P, U, M, or G Mechanism	Yes	501 (53.6)	808 (57.2)
Training Grant	Yes	169 (18.1)	286 (20.2)

Table 5: Characteristics of Articles and Associated Grants According to Whether There Was Sample Size Documentation

Characteristic	Sample Size Estimation		No Sample Size Estimation
Total <i>N</i> (%)		258 (11.0)	2,090 (89.0)
Relative Citation Ratio	Median (IQR)	3.6 (2.1 to 6.4)	2.8 (1.7 to 5.3)
Higher Influence	Yes	152 (58.9)	990 (47.4)
Approximate Potential to Translate	Median (IQR)	0.8 (0.5 to 0.8)	0.5 (0.2 to 0.8)
Higher Translatability	Yes	153 (59.3)	973 (46.6)
Intervention Journal	Yes	239 (92.6)	1,308 (62.6)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	204 (79.1)	878 (42.0)
Number of Grants	Median (IQR)	4.0 (2.0 to 6.0)	3.0 (2.0 to 5.0)
Grant-Years of Funding	Median (IQR)	32.0 (17.0 to 58.0)	24.0 (11.0 to 44.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	28.2 (7.2 to 80.3)	10.8 (3.3 to 35.2)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	2.8 (0.8 to 8.1)	1.2 (0.4 to 4.4)
P, U, M, or G Mechanism	Yes	171 (66.3)	1,138 (54.4)
Training Grant	Yes	66 (25.6)	389 (18.6)

Table 6: Characteristics of Articles and Associated Grants According to Potential to Translate

Characteristic	Higher Translatability		Lower Translatability
Total <i>N</i> (%)		1,126 (48.0)	1,222 (52.0)
Relative Citation Ratio	Median (IQR)	5.1 (3.2 to 8.1)	2.0 (1.3 to 2.9)
Higher Influence	Yes	866 (76.9)	276 (22.6)
Randomization	Yes	378 (33.6)	351 (28.7)
Blinding	Yes	443 (39.3)	492 (40.3)
Sample Size Estimation	Yes	153 (13.6)	105 (8.6)
More Rigorous	Yes	634 (56.3)	634 (51.9)
Intervention Journal	Yes	836 (74.2)	711 (58.2)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	695 (61.7)	387 (31.7)
Number of Grants	Median (IQR)	4.0 (2.0 to 6.0)	3.0 (2.0 to 5.0)
Grant-Years of Funding	Median (IQR)	27.0 (12.0 to 50.0)	23.0 (10.0 to 41.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	16.1 (4.1 to 51.1)	9.6 (3.1 to 29.8)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	1.9 (0.6 to 6.1)	1.1 (0.4 to 3.9)
P, U, M, or G Mechanism	Yes	674 (59.9)	635 (52.0)
Training Grant	Yes	242 (21.5)	213 (17.4)

Table 7: Characteristics of Articles and Associated Grants According to Level of Influence

Characteristic		Higher Influence	Lower Influence
Total N (%)		1,142 (48.6)	1,206 (51.4)
Relative Citation Ratio	Median (IQR)	5.5 (4.0 to 8.3)	1.8 (1.2 to 2.3)
Higher Translatability	Yes	866 (75.8)	260 (21.6)
Randomization	Yes	356 (31.2)	373 (30.9)
Blinding	Yes	438 (38.4)	497 (41.2)
Sample Size Estimation	Yes	152 (13.3)	106 (8.8)
More Rigorous	Yes	614 (53.8)	654 (54.2)
Intervention Journal	Yes	842 (73.7)	705 (58.5)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	733 (64.2)	349 (28.9)
Number of Grants	Median (IQR)	4.0 (2.0 to 6.0)	3.0 (2.0 to 5.0)
Grant-Years of Funding	Median (IQR)	28.0 (12.0 to 52.8)	23.0 (10.0 to 39.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	17.7 (4.3 to 52.5)	8.8 (3.0 to 28.5)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	2.3 (0.6 to 6.4)	1.0 (0.4 to 3.6)
P, U, M, or G Mechanism	Yes	684 (59.9)	625 (51.8)
Training Grant	Yes	261 (22.9)	194 (16.1)

Table 8: Characteristics of Papers and Associated Grants According to Sex of Animals

Characteristic		Male	Female	Both	Unclear
Total N (%)		962 (41.0)	575 (24.5)	474 (20.2)	337 (14.4)
Relative Citation Ratio	Median (IQR)	2.6 (1.7 to 4.5)	2.9 (1.7 to 5.4)	3.3 (1.9 to 6.2)	3.8 (2.0 to 7.1)
Higher Influence	Yes	406 (42.2)	273 (47.5)	265 (55.9)	198 (58.8)
Approximate Potential to Translate	Median (IQR)	0.5 (0.2 to 0.8)	0.5 (0.2 to 0.8)	0.8 (0.5 to 0.8)	0.8 (0.5 to 0.8)
Higher Translatability	Yes	420 (43.7)	268 (46.6)	243 (51.3)	195 (57.9)
Intervention Journal	Yes	613 (63.7)	333 (57.9)	334 (70.5)	267 (79.2)
<i>Nat Med</i> or <i>Sci Transl Med</i>	Yes	269 (28.0)	273 (47.5)	297 (62.7)	243 (72.1)
More Rigorous	Yes	549 (57.1)	238 (41.4)	283 (59.7)	198 (58.8)
Number of Grants	Median (IQR)	3.0 (2.0 to 5.0)	4.0 (2.0 to 6.0)	4.0 (2.0 to 6.0)	4.0 (2.0 to 6.0)
Grant-Years of Funding	Median (IQR)	22.0 (10.0 to 40.0)	27.0 (11.0 to 48.0)	27.0 (11.2 to 48.8)	28.0 (12.0 to 46.0)
Total Grant Funding (2019 \$, Million)	Median (IQR)	8.9 (2.9 to 25.5)	12.8 (4.0 to 39.8)	19.5 (5.2 to 52.2)	17.0 (3.7 to 65.4)
Most Recent Grant Funding (2019 \$, Million)	Median (IQR)	1.0 (0.4 to 3.8)	1.5 (0.5 to 5.1)	2.5 (0.6 to 6.3)	1.8 (0.5 to 7.5)
P, U, M, or G Mechanism	Yes	489 (50.8)	315 (54.8)	309 (65.2)	196 (58.2)
Training Grant	Yes	152 (15.8)	135 (23.5)	97 (20.5)	71 (21.1)

Table 9: Exploratory Logistic Regression Models for Measures of Rigor

	Dependent variable			
	Randomized (1)	Blinded (2)	Sample Size Estimation (3)	More Rigours (4)
Number of Grants	0.095** (0.041)	0.008 (0.033)	0.231** (0.113)	0.037 (0.031)
Grant Years	-0.053 (0.189)	0.011 (0.197)	0.268 (0.292)	-0.052 (0.175)
Most Recent Funding	0.420*** (0.114)	-0.244** (0.105)	0.576*** (0.172)	0.122 (0.103)
U, M, G, or P Grant	-0.132 (0.130)	0.004 (0.120)	-0.156 (0.200)	-0.074 (0.118)
Intercept	-0.591*** (0.186)	-0.529*** (0.177)	-2.500*** (0.298)	0.190 (0.173)
Observation	2,348	2,348	2,348	2,348
R2	0.011	0.005	0.03	0.002
$\chi^2$ (df=5)	17.660***	8.813*	35.902***	2.9

Note: \* $p < 0.1$ ; \*\* $p < 0.05$ ; \*\*\* $p < 0.01$

### Recommendations

The subcommittee identified a number of possible financial ramifications of changes to the way in which NIH funds and oversees animal research. Informed by those possibilities, we linked methodological rigor data from [Ramirez et al.](#) with data on article metrics and grant history. Exploration of these descriptive data suggest that higher levels of rigor do not necessarily imply higher costs, with the exception of sample size estimation. Higher levels of translatability may be associated with higher levels of rigor and higher costs. The subcommittee offers the following recommendations:

- NIH should develop an evaluation program to assess the progress in implementing recommendations, their effects on both NIH and the external research community, and challenges that arise in implementing recommendations.
- NIH should conduct and support analyses on the elements and impact of rigor and transparency in grant applications and publications to better understand the impact of measurable factors on quality, cost, and rigor in NIH-funded research.
- NIH should allow applicants to include text in the budget justification section on how projected animal budgets are linked to efforts to enhance transparency, rigor, and translatability.
- NIH and/or other analysts should consider studies of scientists who demonstrate the highest levels of transparency and rigor; they may offer lessons that could be applied enterprise-wide.

## Appendix 6 — Example of Effective Practices to Enhance Rigor and Transparency

The following power calculation was an example provided by Catherine Kaczorowski, Ph.D., whose research at The Jackson Laboratory focuses on a collaborative and interinstitutional research program that uses a multidisciplinary approach to identify early causative events underlying “normal” nonpathological age-related memory decline and Alzheimer’s dementia. As part of a presentation we saw examples of effective practices they have adopted.



### Power calculation for causal test via CRISPR genome editing (R markdown file)

#### ApoE x Tomm40 - Power analyses and future plans

David J Anderson September 23, 2019

##### Introduction: What is this analysis?

Before we conduct statistical tests for effects of Genotype, age, and sex on CRISPR mouse CFA/CFM, we need to make sure these tests are well-powered. Really, this should be done at the outset of experiments to determine how many mice we need to breed, phenotype and harvest, but as far as I can tell only rule-of-thumb estimates were used. This is understandable as I myself have had a fair bit of difficulty tracking down the requisite information to run this analysis (namely, expected effect sizes for each variable), but the following summarizes my attempts to figure out what sample sizes we’ll need to feel confident moving forward with our analysis. I begin with my calculations of effect sizes for genotype, age, and sex on CFA and CFM tasks based on Neuner’s papers. I then calculate the sample size necessary for each group for our analyses to be well-powered. Finally, I summarize these data and compare the sample sizes required to the number of datapoints we already have and the number of mice we have coming through the pipeline currently. This should serve as a roadmap for what mice to breed and phenotype going forward.

**Calculation of effect sizes:** I calculated expected effect sizes for Genotype, Age, and Sex from the F-ratios and N’s reported in [Neuner’s Neuron paper](#) using the calculators available [here](#). Across AD-BXD mice, there was a significant effect of Apoe allele ( $F(1,354) = 4.7$ ;  $p = 0.03$ ), age ( $F(1,354) = 12.3$ ;  $p = 0.001$ ), and sex ( $F(1,354) = 17.9$ ;  $p < 0.001$ ) on CFA.

IV	Variable	F statistic	Effect Size (d)
Genotype	CFA	4.7	0.226/0.85*
Age	CFA	12.3	0.283
Sex	CFA	17.9	0.348
Genotype	CFM	20.9	0.476/1.79*
Age	CFM	86.2	0.75
Sex	CFM	4.9	0.182

**Calculating group size:** From the table, we can see that the effect sizes of genotype and age are smaller for CFA while the effect size for sex is smaller for CFM. As CFM/CFA tests are paired (i.e. any mouse that gets scored for one will be scored for the other) and as we want to be able to test for differences in both variables I feel that it is fair to use the *smaller* effect sizes for each variable when calculating desired group sizes. Let me know if this doesn’t make sense; I’m not sure if I’m explaining my reasoning well. Running power analysis for genotype and age is easy enough if we just assume simple ANOVA models; there are ways to do this for more complex models, but I’m not well versed in them: this might be something we should talk about going forward.

**Genotype Power Analysis:** To start with, I calculated group sizes per genotype with  $d = 0.226$ . I’m assuming the number of groups ( $k$ ) to be 12; we have *many* genotypes currently, and I’m assuming we’ll ultimately want to look for differences between all of them.

Balanced **one-way** analysis of variance power calculation `pwr.anova.test(k=12, f=0.226, power=0.8)`

k = 12

n = 28.24219

f = 0.226, sig.level = 0.05, power = 0.8

This calls for ~28 mice to be phenotyped in each genotype  
Bred, enter pipeline (all blind until final data for all QCed)