



COSMETIC SURGERY  
FOUNDATION

# Cosmetic Surgery Foundation

## 2026 Research Grants

### Instructions & Guidelines

**Submission Deadline: October 6, 2025**

The Cosmetic Surgery Foundation (CSF) supports the research and educational programs of the American Academy of Cosmetic Surgery and builds relationships that advance the science and safety of cosmetic surgery through innovative, targeted funding and inspired leadership.

#### General Information:

The research grants are in the amount of \$2,000. All monies are to be spent and the project completed within 12 months. Half of the grant money is provided in December. The other half is provided in June 2026, but only after receiving the mid-term progress report. Final reports are due November 14, 2026. Grant recipients are expected to submit an abstract of their study for the 2026 AACS Annual Scientific Meeting. Grant recipients are also expected to submit a manuscript of their study for publication in the *American Journal of Cosmetic Surgery*.

All research grant applications are rated and selected by the CSF Board of Directors. Grant recipients are recognized with certificates at the 2026 AACS Annual Business Meeting at the Annual Scientific Meeting.

#### Eligible Candidates Must Be:

- An AACS member in good standing, **and**
- A physician in the United States (MD, DO, DDS, DMD), **and**
- A Resident or Training Fellow

#### Timeline:

Submission Deadline:	October 6, 2025, midnight
Rating:	October 7-13, 2025
Decisions Announced:	November 8, 2025
1 <sup>st</sup> Checks (50% of amount):	December 10, 2025
Certificates Presented:	February 6, 2026 (in Tampa)
Mid-Term Reports Due:	May 1, 2026
2 <sup>nd</sup> Checks (50% remaining):	June, 2026
Abstracts for ASM Due:	May-June 2026
Final Reports Due:	November 14, 2026

#### How to Apply:

By September October 6, 2025, submit all application materials via the online form at: <https://foundation.cosmeticsurgery.org/grant-applications/>

(Continued on next page)



# CSF 2026 Research Grants

**Submission Deadline October 6, 2025**

**Information requested is as follows:**

- 1) General information (name, address, etc.)
- 2) Curriculum Vitae
- 3) Headshot photo for recognition
- 4) Research grant proposal – One document that includes the following:
  1. **Title of Research Project**
  2. **Lead Researcher**
  3. **Other Research Team Members, if applicable**
  4. **Amount Requested**
  5. **Project Aims:** What do you intend to accomplish? What hypothesis is to be tested?
  6. **Importance:** Why is the research important? Evaluate existing knowledge in the field and specifically identify the possible contributions that your investigation may make.
  7. **Preliminary Studies:** What has already been done in this field?
  8. **Experimental Design:** How are you going to accomplish the research? Describe the experimental design, the procedures to be used, and the manner in which the data will be analyzed. Do not include established laboratory procedures.
  9. **Literature Cited**
  10. **IRB or Ethics Committee Approval:** Indicate if your study has been reviewed and approved by an IRB or Ethics Committee. If so, include approval letter. (See attached article for more information about IRBs.)
  11. **Background:** Please describe the organizational structure in which the research will be carried out and include a supporting letter from the sponsor/preceptor/program director describing the nature of the linkage to a division or department. Include professional background and/or academic affiliation.
  12. **Disclosures:** If applicable, include any relevant financial disclosures.
  13. **Budget:** Create a line-item budget specific to the amount of funding you are requesting. The CSF does not fund: salaries, travel, lodging, publishing fees, overhead costs, or items such as ipods, laptops, cameras, etc.
- 5) Signed Research Grant Terms & Conditions

**For Questions:**

Please feel free to reach out to us with questions.

Cosmetic Surgery Foundation  
1932 S. Halsted Street, Suite 413, Chicago, IL 60608, USA  
Phone: 1-312-981-6760  
[info@cosmeticsurgery.org](mailto:info@cosmeticsurgery.org)

# The American Journal of Cosmetic Surgery

## The Institutional Review Board (IRB): What It Is and Why You Need One

Jane A. Petro, MD, FACS, FAACS, Editor Emeritus



The Institutional Review Board, or IRB for short, is also known as an independent ethics committee (IEC), ethical review board (ERB), or research ethics board (REB). It generally consists of a formally designated committee that reviews and monitors research projects that intend to use human beings as the research objects. For a comprehensive look at IRBs in question and answer

format, please [click here](#). The IRB is in place to protect the rights and well being of human research participants from both physical and psychological harm. It also serves to assure that whatever harm involved is explicitly described and that the subjects 1) understand the risks, and 2) agree to participate of their own volition. The IRB may be sponsored within the institution conducting the research or it may be an independent entity tasked with performing IRB functions. If you are interested in conducting prospective clinical research, IRB approval is required and must be obtained from your university or hospital IRB or a commercial entity.

IRBs were created in response to explicit harms that came to participants in research done in the United States (the infamous [Tuskegee Syphilis Study](#), for example, or radiation experiments done by the military) and in Europe as part of Nazi “research.” The [Belmont Report](#) identified three key ethical principles of human research: 1) respect for persons, 2) beneficence, and 3) justice. An IRB may only approve research for which the risks to subjects are balanced by potential benefits to society, and for which the selection of subjects presents a fair or just distribution of risks and benefits to eligible participants. For more information on the IRB, [click here](#).

Even retrospective studies, if involving humans, as is common in retrospective case reviews, may benefit from an IRB-approved exemption. Research involving the analysis of existing data and other materials if they are already publicly available, or where the data can be collected such that individual subjects cannot be identified in any way, does not require IRB approval. [Click here](#) for a detailed overview of these issues (Parker GE. A framework for navigating Institutional Review Board (IRB) oversight in the complicated zone of research. *Cureus*. 2016; 8(10):e844. [doi:10.7759/cureus.844](#)). Commercial IRBs are acceptable alternatives to standard hospital- or university-based review boards.

It is good and established practice for medical journals to avoid publishing research that has not met IRB standards. The following requirements are important to keep in mind if you are considering research that involves human participants. You must

- Obtain prior approval for human subjects research by an IRB or equivalent ethics committee(s).

- Declare compliance with ethical practices upon submission of a manuscript.
- Report details on how informed consent for the research was obtained (or explain why consent was not obtained).
- Submit, upon request from the journal, documentation from the review board or ethics committee confirming approval of the research.
- For clinical trials, provide trial registration details, the study protocol, and [CONSORT](#) documentation (which is a standard evidence-based, minimum set of recommendations for reporting randomized trials; for more information, [click here](#)).
- Confirm that an identified individual has provided written consent for the use of that information.

### Member Resource: A Commercial IRB

#### Pearl IRB

29 East McCarty Street, Suite 100  
Indianapolis, IN 46225  
[www.pearlirb.com/](http://www.pearlirb.com/)

Pearl IRB is an independent institutional review board, fully accredited by the Association for the Accreditation of Human Research Protection Program Inc. (AAHRPP). Pearl IRB manages the local and central IRB needs for large and small institutions, principal investigators, CROs, and sponsors. Its vision is to improve the clinical research process which will lead to delivering therapeutics and diagnostics to patients sooner. Whether the research is a small single site project or a complex multi-center study, Pearl IRB can help. If you are interested, contact Pearl IRB for its fee schedule.

If you're unsure how your study may be classified for review, the [Office for Human Research Protections \(OHRP\)](#) has some helpful resources. [Click here](#) to review their [online decision trees](#). These charts are designed to help you decide if an activity is research involving human subjects that must be reviewed by an IRB and whether informed consent, or documentation of informed consent, can be waived.

Pearl IRB board review turnaround times are 1-3 business days for exempt or expedited reviews, and 8-10 business days for a full board review.