Guidelines
Clinical Research Units
1 Programme Information

1 Objective

Funding for a Research Unit enables researchers to collaborate closely on specific medium-term projects whose anticipated findings could not be achieved within the scope of the Research Grants Programme. A Research Unit typically has fewer than ten projects which are coordinated to enable work on a common research topic. The goal of a Clinical Research Unit is to promote research networks whose thematic focus is on translational clinical research with patient-oriented approaches and on understanding disease mechanisms. All other research topics are covered by the “Guidelines for Research Units” (DFG form 50.04).

In addition to meeting the criteria that apply to all Research Units, Clinical Research Units are also expected to enhance the scientific profile of their main location and institution, and improve the quality of clinical research by creating and strengthening research-oriented structures within university hospitals. In addition, they should increase cooperation between clinicians and scientists in the foundational disciplines of medicine, improve training structures for clinical research, and support the performance-based allocation of resources.

2 Proposals

2.1 Eligibility

A proposal for a Clinical Research Unit is prepared jointly by several researchers and submitted by the spokesperson. Researchers must work in Germany and have completed their academic training (usually with a doctorate).

Researchers who work at an institution that is not non-profit, or one that does not allow immediate publication of research findings in a generally accessible form, are not eligible to apply.

A Clinical Research Unit is primarily run by university hospitals at one location. Cooperation with non-university research institutions is possible if the majority of research projects are based at universities. The involvement of external scientists at other locations or abroad is only possible in justified exceptional cases.
2.2 Format and deadline

Applicants must first submit a draft proposal to the DFG Head Office, which is then forwarded to reviewers. If successful, the DFG will invite the applicants to submit a full proposal. Draft proposals and full proposals may be submitted at any time. For more detailed information on preparing your proposal, please consult the instructions for proposals to establish Clinical Research Units.

[www.dfg.de/formulare/54_09/](www.dfg.de/formulare/54_09/)

Proposals for the individual projects within the Clinical Research Unit should be submitted according to the instructions for project proposals.

[www.dfg.de/formulare/54_01/](www.dfg.de/formulare/54_01/)

3 Duration

The total funding duration is generally six years, or eight years in exceptional cases. The first funding period is three years. Further funding requires the submission and approval of renewal proposals.

4 Participants

The Clinical Research Unit consists of the leaders of the individual projects. One of the researchers assumes the role of the spokesperson, who represents the Clinical Research Unit toward the DFG and third parties, and is required to submit reports to the DFG. The spokesperson should be a full-time university teacher.

In addition to the spokesperson, the Clinical Research Unit will also be headed by a research coordinator. Once the Clinical Research Unit is established, the person appointed to the research professorship assumes its scientific and administrative leadership as research coordinator. The person appointed to the research professorship must meet particular requirements with regard to his/her scientific track record, experience in leading projects (including projects with third-party funding), and integration and leadership skills. These criteria will be considered during the review process. The spokesperson or the research coordinator is responsible for managing the unit's common central-project funds, especially those for coordination.
In exceptional cases, the spokesperson of the Clinical Research Unit may also serve as its research coordinator and hold the research professorship.

II Proposal Modules

To apply for funding in the Clinical Research Units Programme, the project leader may submit the modules listed below for the individual research projects. Please see the respective guidelines for additional information on the modules.

1 Basic Module

Use the basic module to request funding for direct project costs, project-specific staff, and instrumentation necessary to carry out the project. In addition to funding for purely scientific projects, Clinical Research Unit proposals may also request funding for pilot studies and other necessary patient-oriented research costs (e.g. specialised outpatient clinics, resources for building and expanding infrastructure for clinical trials, archiving of patient data).

www.dfg.de/formulare/52_01/

2 Replacements

If your project requires that you be released from teaching or administrative duties, you can use this module to request funding for a replacement to take over these responsibilities.

www.dfg.de/formulare/52_03/

The following modules can be submitted by the project leaders for the individual projects in conjunction with the modules above, or by the spokesperson or research coordinator for the entire Clinical Research Unit within the coordination proposal:

3 Temporary Substitutes for Clinicians

If this project requires that clinicians conduct research, you can use this module to request funding for temporary substitutes to take over their patient-care responsibilities.

Each Clinical Research Unit must apply for lump-sum funding of at least one temporary substitute. The department of medicine must finance the funding for this position.
(from its state appropriations for research and education). The DFG grants lump-sum funding for one additional temporary substitute position (or 50% of each position if funding is requested for multiple positions). Furthermore, the DFG makes available, upon request, flexible funds in the amount of €60,000 per year to support especially creative structural measures in hospitals and clinics.

www.dfg.de/formulare/52_04/

4 Project-Specific Workshops

If you would like to conduct workshops for your projects or for the entire Clinical Research Unit, you may request funding to help you do so.

Please use this module to request funding for all workshops and colloquia that you will conduct within the funding period or for a concluding colloquium. It is not possible to submit a request for colloquium funds at a later date.

www.dfg.de/formulare/52_06/

5 Mercator Fellows

This module enables you to pursue an intensive and long-term exchange with researchers in Germany and abroad. Fellows will partially be on site but will remain in contact with the Clinical Research Unit even after their stay.

www.dfg.de/formulare/52_05/

6 Public Relations

To enable you to present your work to the general lay public, you can request funding for public relations.

www.dfg.de/formulare/52_07/

The following modules can only be submitted by the spokesperson or the research coordinator on behalf of the entire Clinical Research Unit within the coordination proposal:
7  Professorship

To support the Clinical Research Unit, a professorship may be established with advance and/or partial funding by the DFG. Such funding must enable an early professorial appointment or the implementation of a structural improvement.

The establishment of a research professorship at the main hospital must be part of the proposal, unless the holder of an existing research professorship will assume the scientific leadership of the Clinical Research Unit. At the time of submitting the proposal, the department of medicine must certify in writing that it will make the research professorship permanent after the fourth year of funding and support it for at least five years after DFG funding expires (with two research assistants and two technical assistants, plus an appropriate annual allowance for consumables in the amount of €50,000).

The scientific topic of the professorship must be in line with the mission of the Clinical Research Unit. The research professor’s main focus must be collaboration within the unit. For this reason, the department of medicine must exempt the research professor, upon the research coordinator’s request, from any clinical obligations.

www.dfg.de/formulare/52_10/

8  Coordination

The spokesperson or the research coordinator may apply for funds needed to coordinate the various projects within the network.

www.dfg.de/formulare/52_12/

9  Network Funds

Use this module to request funds for the entire network.

www.dfg.de/formulare/52_13/

10  Start-Up Funding

Through this module, research networks can receive funds to help promising early career researchers pursue independent projects.

www.dfg.de/formulare/52_11/
11 Gender Equality Measures in Research Networks

This module enables research networks to implement targeted measures to promote gender equality in science and academia. For this purpose, up to €50,000 of funding may be requested per three-year funding period. These funds may also be used to enable research physicians and other scientists who return from parental leave to lead projects and to submit regular funding proposals for projects within the Clinical Research Unit.

www.dfg.de/formulare/52_14/

III Special Provisions

1 Co-financing and budgetary commitment

Funding for Clinical Research Units requires co-financing from the state appropriations for research and education to the university hospitals and medical departments, as well as the allocation of performance-based funding to the applicants in addition to core support. Co-financing must ensure that:

- after three years of DFG support for the proposed research professorship and a positive interim evaluation, from the fourth year on, the research professorship will be funded in full by the university’s department of medicine from the state appropriations for research and education, and will be included in the regular hospital/medical department budget for at least five years after DFG funding for the Clinical Research Unit expires (usually after six years), along with the necessary core support to ensure scientific viability (two research assistants and two technical assistants, plus an appropriate annual allowance for consumables in the amount of €50,000);

- at least one temporary substitute position in the Clinical Research Unit (and, if more than one such position is proposed, 50% of all such positions) will be financed from the state appropriations for research and education. If one temporary substitute position is approved, the DFG will fund an additional one. Furthermore, the DFG makes available, upon request, flexible funds in the amount of €60,000 per year to support especially creative structural measures in university hospitals and clinics.
No co-financing is expected for any projects outside of the department of medicine.

If funding is approved, the DFG pays out its share of the overall budget for the Clinical Research Unit. Accounting for the unit is expected to also identify funds derived from state appropriations and how they were used in the project. To simplify accounting, expenditure reports must include both those expenses that were covered by the DFG and the overall costs of the Clinical Research Unit (divided into payroll and direct project expenses).

2 Systematisation and structuring of doctoral training for medical researchers

The proposal must include a statement by the department of medicine describing any plans to systematise and structure doctoral training for medical researchers or to structure scientific training. Doctoral positions for medical researchers may be proposed in addition to standard staff funding (grants for research sabbaticals). They must be funded by the department of medicine following a favourable evaluation.

3 Association of Emmy Noether Independent Junior Research Groups

Clinical Research Units may associate Emmy Noether Independent Junior Research Groups that investigate related topics. In this case, the Clinical Research Unit proposal and the Emmy Noether proposal should refer to each other. Decisions on both proposals will be made independently of each other. If both the Clinical Research Unit and the Emmy Noether group are established, the junior research group leader will participate in the Clinical Research Unit’s shared events. The association can also be established afterwards at the spokesperson’s discretion.

4 Transfer projects

Transfer projects are based on results generated by DFG-funded research projects. They serve to test scientific insights in practice and, in collaboration with an application partner, develop basic-research findings into prototypes or exemplary applications. The application partner may be a commercial enterprise or a non-commercial, non-profit institution, especially in the public sector.

The core of the project is a joint work programme, focusing on intensive mutual exchange of scientific knowledge and corresponding application issues.
Further information on transfer projects can be found in the relevant proposal instructions.

www.dfg.de/formulare/54_014/

IV Obligations

In submitting your draft or full proposal to the DFG, you agree to:

1. adhere to the rules of good scientific practice.¹

The general principles of good scientific practice include, among others: maintaining professional standards, documenting results, rigorously questioning all findings, and attributing honestly any contributions by partners, competitors and predecessors.

Scientific misconduct is defined as the intentional and grossly negligent statement of falsehoods in a scientific context, the violation of intellectual property rights or impeding another person’s research work. The circumstances of each case will be considered on an individual basis. In cases where scientific misconduct has been established, the DFG may impose one or more of the following sanctions, depending on the nature and severity of the scientific misconduct:

- issuing a written reprimand to those involved;
- exclusion from the right to apply for DFG funds for a period of one to eight years, depending on the severity of the scientific misconduct;
- revoking funding decisions (complete or partial cancellation of the grant, recalling granted funds, demanding repayment of funds spent);

¹ The rules of good scientific practice are presented in detail in the white paper entitled “Safeguarding Good Scientific Practice” and in the usage guidelines for research grants (DFG form 2.01).
demanding that those concerned either retract the discredited publications or correct the falsified data (in particular by publishing an erratum), or appropriately indicate the DFG’s retraction of funding in the discredited publications;

- exclusion from acting as a reviewer or from membership in DFG committees for a period of one to eight years, depending on the severity of the scientific misconduct;

- denying voting rights and eligibility in elections for DFG statutory bodies and committees for a period of one to eight years, depending on the severity of the scientific misconduct.

By accepting funding, the recipient agrees to

2. use the grant exclusively and in a targeted manner to realise the funded project. The use and accounting of funds must conform to the relevant regulations of the DFG.

3. submit progress reports on the research according to the dates specified in the award letter and to present financial accounts to the DFG detailing the use of funds.

The DFG expects that the findings of the projects it funds be made available to the public.

V  Publication of Data on Grant Holders and Research Projects

The data necessary for processing your grant proposal will be stored and processed electronically by the DFG. If a grant is awarded, your work address (e.g. telephone, fax, e-mail, internet website), as well as information on the content of your research project (e.g. topic, summary, keywords, international cooperation), will be published in the DFG’s project database GEPRIS and – in excerpts (grant holder’s name, institution and location) – in the “Programmes and Projects” section of the DFG’s electronic annual report. If you do not wish this information to be published electronically, please notify us in writing no later than four weeks after receipt of your award letter.

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