





Research Nurse for the Clinical Research Support Unit

What do we offer?

Scientific-Technical Service	Scientific-Technical Service Clinical Research Support Unit
	(USIC)
Type of contract	Indefinite with a 6-month probationary period
Beginning	Immediate
Day	37.5h/week – full-time
Category	Senior Research Technician C4
Remuneration	€31,294.37 gross/year

The Institute

The Biomedical Research Institute of Lleida, Dr. Pifarré Foundation (IRBLleida) was created with the aim of creating synergies between basic, clinical and epidemiological research, making biomedical research a key driver to improve current clinical practice. IRBLleida covers a chain of translational research, from basic research, aimed at understanding the physiological and pathological mechanisms of the human body, to research that studies the behavior of diseases in large population groups.

IRBLleida was founded in 2004 through a collaboration agreement between the University of Lleida (UdL) and the Catalan health system. IRBLleida integrates research groups from the faculties of Medicine and Nursing and Physiotherapy of the UdL. On the other hand, we incorporate research groups from:

1. The Catalan Institute of Health (ICS) both in the hospital field (Hospital Universitari Arnau





de Vilanova-HUAV) and in primary care in Lleida and the Alt Pirineu-Aran Health Region.

2. The healthcare provider Healthcare Service Management (GSS; Santa Maria-HUSM University Hospital, Pallars Regional Hospital and Mental Health, among others).



IRBLIeida has been a CERCA institute since 2013, and consequently it is organized according to a model of good governance and operation that guarantees efficiency, flexibility of management, recruitment and promotion of talent, strategic planning and executive capacity.

In addition, it is one of the 34 Spanish Health Research Institutes (IIS) recognised by the Carlos III Health Institute and the Government of the Generalitat, as established by Law 16/2003, of 28 May, on the cohesion and quality of the national health system.



In December 2014, the Biomedical Research Institute of Lleida received the 'HR Excellence in Research' award from the European Commission. It is a HR EXCELLENCE IN RESEARCH recognition of the Institute's commitment to developing a human resources strategy for researchers, designed to align practices and procedures with the principles of the <u>European Charter for Researchers</u> and the <u>Code of Conduct</u> for the recruitment of researchers.

Check out our recruitment policy.





Professional profile of the person hired

Research Nurse

Requirements (those candidacies that do not meet this point will be excluded)

- Bachelor's degree in nursing or equivalent
- Professional experience in nursing

These requirements must be met at the beginning of the contract.

Tasks to be carried out

The selected candidate will join the Scientific-Technical Service (SCT) Clinical Research Support Unit (USIC) of IRBLleida and will provide support by carrying out the following tasks:

- Control and management of nursing procedures and complementary tests according to each protocol.
- Comprehensive support for patient care.
- Patient and test scheduling.
- Pharmacovigilance: management of adverse events.
- Processing of biological samples
- Clinical trial coordination functions:
 - o Participate in the selection of the centre and the research team.
 - Participate in the evaluation of the feasibility of the study, analysis of the circuits and technical needs, spaces and personnel of each clinical trial.
 - Conducting meetings with researchers and facilitating communication between the medical services involved in the study and external stakeholders.
 - o Maintenance of necessary files: researcher, promoter, pharmacy.
 - o Facilitate the process of recruitment and obtaining informed consent.





- Coordination of visits and follow-up controls.
- Management of possible audits.
- Contact with monitors and preparation of documentation for external monitoring.
- Recording of the data in the data collection notebook, resolution of queries and monitoring.

It will be valued

Knowledge

- Master's Degree in Clinical Trials or Clinical Research
- Training related to the field of Clinical Trials
- Current Certificate of Good Clinical Practices

Experience

- Previous experience in a similar position within the Clinical Trials sector or in the area of clinical research
- Experience with handling electronic data collection notebooks
- Experience with handling SAP software
- Experience in the processing of biological samples

Competences

- Organisational capacity
- Teamwork
- Proactive attitude





Characteristics of the contract

- ✓ Permanent contract with a six-month probationary period
- ✓ Full-time (37.5h/week)
- ✓ Immediate start

Why work at IRBLleida?



We offer a highly stimulating environment with state-of-the-art infrastructures.



We offer complementary training for all profiles. To view our training and development portfolio, please visit our website in the training section <u>.</u>



We offer and promote a diverse and inclusive environment and welcome applicants regardless of age, disability, gender, nationality, race, religion or sexual orientation.



Work-life balance and the possibility of benefiting from flexible working hours. In addition, as a result of different company agreements, the following improvements are recognized:

- Paid leave to go to the doctor for personal health reasons.
- Paid leave to accompany a first-degree family member under 18 years of age, over 70 years of age or with a first degree of disability to the doctor.
- Public holidays that coincide with Saturday or Sunday are moved to the Monday immediately following.
- A special 6-hour day is established on Holy Thursday, April 23, June 23,
 December 24, December 31 and January 5.





Documentation and deadline for submission

Applications must be accompanied by:

- Cover letter.
- Resume.

The deadline for submission will end on September 24, 2025 at 2:00 p.m.

Applications received after the deadline/time will be automatically excluded.

Interested persons can apply for the offer by filling in the <u>form</u> and sending their CV and cover letter, indicating the name of the offer to which you are applying and the reference **051-25**.

Selection calendar for the reference process 051-25			
Minimum 15 days	Publication and dissemination of the offer: IRBLleida		
	website, Euraxess (for research staff), social networks,		
	other employment websites depending on the position		
	offered.		
Maximum following 2 working days	Sending CVs to the Selection Committee		
	Celebration of the Selection Committee		
Maximum following 5 working days	- Interview with the pre-selected candidates		
Waxiii alii Tollowiiig S Workiiig days	- Assessment and Award Minutes of the		
	Selection Committee		
Maximum following 5 working days	Carrying out the administrative procedures necessary		
	to formalise the employment contract		
	Immediate		
Approximate start of the contract			





Express selection process

In those cases in which a worker has to be replaced urgently, for example, to cover sick leave, because for scientific reasons the incorporation must take place on a specific day, because it is provided for in a resolution, etc., an express selection procedure may be followed.

This selection process will follow the same procedure as the ordinary one, but the duration of all phases of the process will be reduced, mainly the phase of publication of the job offer and submission of applications and the phase of evaluation and selection of personnel.

Regulation and regulatory principles

The hiring will be carried out in accordance with the provisions of art. 15 of Royal Legislative Decree 1/1995, of 24 March, approving the text of the Workers' Statute Law, in accordance with the provisions of art. 2 of Royal Decree 2720/98, of 18 December (B.O.E. of 8 January 1999), Law 12/2001, of 9 July (B.O.E. of 10 July) and concordant provisions.

The principle of equality between men and women is taken into account, in accordance with **Organic Law 3/2007, of 22 March**, for the effective equality of women and men. IRBLleida has an <u>Equal Opportunities Plan for men and women</u> and a <u>Protocol for the prevention and eradication of sexual harassment.</u>

The right to equal opportunities and treatment is taken into account, as well as the real and effective exercise of rights by people with disabilities on equal terms with other citizens, through the promotion of personal autonomy, universal accessibility, access to jobs, etc. inclusion in the community and independent life and the eradication of any form of discrimination, in accordance with articles 9.2, 10, 14 and 49 of the Spanish Constitution and the International Convention on the Rights of Persons with Disabilities and the international treaties and agreements ratified by Spain, in accordance with the provisions of Royal Legislative Decree 1/2013, of 29 November.





Reservation of places for people with disabilities

In accordance with the provisions of Article 42 of Royal Legislative Decree 1/2013, of 29 November, approving the Revised Text of the General Law on the Rights of Persons with Disabilities and their Social Inclusion, this call reserves a **percentage of no less than 2%** of the places to be covered by people who can prove a disability equal to or greater than 33%.

Applicants who wish to opt for this reservation must submit documentation accrediting their disability and, where appropriate, request the necessary adaptations for the selection tests.

In the event that the reserved places are not covered due to a lack of applicants who meet the requirements, these will be accumulated in the general access places.

**The text of this document has been written in Catalan, Spanish and English, considering the three versions as official, but in case of conflict the Catalan version will prevail.

IRBLleida is committed to the principles of Merit-Based Recruitment and Transparency

(OTM-R) in accordance with the HRS4R seal requirements





ANNEX I. MEMBERS OF THE SELECTION COMMITTEE

President	Ms. Eva López, Manager of IRBLleida
Vocal	Dr. Alicia Sánchez, Head of the Clinical Research Support Unit
	Dr. María Ruiz, coordinator of the SCT of IRBLleida
Secretary	Ms. Elena Moscatel, Head of the Department of People and Legal





ANNEX II. SCALE OF MERITS

Academic curriculum and complementary training	30 Points
Master's Degree in Clinical Trials or Clinical Research	10 points
Training related to the field of Clinical Trials	15 points
Current Certificate of Good Cynical Practices	5 points
Accredited professional experience	50 Points
Previous experience in a similar position within the Clinical	20 points
Trials sector or in the area of clinical research	
Experience with handling electronic data collection notebooks	10 points
Experience with handling SAP	10 points
Experience in the processing of biological samples	10 points
Competency test or interview	20 points
Criteria subject to value judgment will be assessed in accordance	20 points
with the interview carried out	
Maximum score	100 points

Applications that do not exceed 50% of the maximum score will be rejected





Data protection information clause

Data controller

Identity: BIOMEDICAL RESEARCH INSTITUTE OF LLEIDA

CIF: G25314394

Postal address: Av. Alcalde Rovira Roure nº80, 25198, Lleida

Email: protecciodedades@irblleida.cat

Purpose of data processing and storage

At **the LLEIDA BIOMEDICAL RESEARCH INSTITUTE** (hereinafter referred to as **IRBLLEIDA**) we process the information you provide us as an interested party, to manage the processing of your CV and candidacy.

The data obtained will be kept for a period of up to 12 months, to cover future applications if they are not updated before or until there is opposition to their processing by the interested party.

Legitimacy for data processing

The legal basis for the processing of your data is the consent of the interested party when contacting the LLEIDA BIOMEDICAL RESEARCH INSTITUTE.

Recipients of your data

Your data will be communicated to third parties and collaborators related to the organization. Apart from these entities, your data will not be communicated to third parties.

Rights of the interested parties

The owners of the data processed by **IRBLLEIDA** have the right at any time to access their data, rectify them, oppose their processing or delete them if they believe that they are no longer necessary for the purposes for which they were collected. In addition, if you wish, you may request the portability of your data and limitation of its processing. In the latter case, we will only keep them for the exercise or defence of claims. You may also revoke the consent granted at

To exercise these rights, you can contact **IRBLLEIDA** through the email <u>protecciodedades@irblleida.cat</u>. Likewise, if you consider that your rights have been violated, you may file a complaint with the Catalan Data Protection Authority.