





| | | JOB OFFER | | |
|---|---|--|---|--|
| REFERENCE SANOFI23/29 | | OPENING DATE | DEADLINE | |
| | | 17/10/2024 | 26/10/2024 | |
| | PRC | FILE REQUIREMENTS | | |
| | EXCL | USIVE REQUIREMENTS: (1) | | |
| ACADEMIC DEGREE | | health sciences or life sicences + of vided with the application). | ficial master/Equivalent (Justification | |
| | | ALUED MERITS /SKYLLS | | |
| FURTHER | Degre Degre Biome Maste Speci certifi datab Know clinica Traini mana | ee in Statistics, Degree in Physics o edical Engineering. r's degree in biomedical research, fic training in database managemen icates in tools such as SPSS, R, STA bases. ledge of research methodology: Sta al study design and scientific writing ng in project coordination or clinical gement and coordination of multice | rmation Sciences (Documentation), r Mathematics or Degree in public health or epidemiology. nt and statistical analysis: Courses o ATA or management of scientific udies or complementary training in g. I studies: Courses or certificates on | |
| EXPERIENCE | health area. Experience in clinical data collection and processing, database management, coordination of multicentre studies, bibliographic research, writing scientific articles, collaboration in research projects, use of statistical analysis tools. Experience in processing administrative documentation for ethics committees and clinical research management Experience in research management | | | |
| LANGUAGES | Englis | sh (Certificates or diplomas must be | e provided). | |
| OTHERS | | Clinical Practice (GCP) accreditation | n. | |
| | CON | TRACT INFORMATION | | |
| TYPE OF CONTRAC | СТ | EXPECTED INCORPORATION DATE | JOB STATUS | |
| Contract for scientific-technical activities (article 23.bis of Law 14/2011, of June 1, on Science, Technology and Innovation) | | October/November | Full time. 1.710 hours per year (approx. 37,5 h/week) | |
| ANNUAL GROSS SAL | ARY | DURATION O | F THE CONTRACT | |
| 28.487,38 € without prejudice update established in state le 2024. | | | ation of the project or to external public grants in full competition). | |
| WORK LOCATIONS | | UNIT/DEPARTMENT | | |
| Hospital Universitario Marqués de Valdecilla | | Otorhinolaryngology / Rhinology Section | | |
| | | JOB DETAILS | | |
| | | OFFER DESCRIPTION | | |
| | Res | search support technician | | |
| | | FUNCTIONS | | |
| the databases associated wi | th the CRAIS | study, ensuring the correct co | e comprehensive management ollection, storage and analysis ing efficient communication an | |

For the position of Research Support Technician, key functions include the comprehensive management of the databases associated with the CRAIS study, ensuring the correct collection, storage and analysis of data. In addition, you will coordinate the participating centres, ensuring efficient communication and compliance with the project objectives. Within the research tasks, an exhaustive bibliographic search will be carried out to keep the relevant scientific references and advances up to date, as well as the writing of research articles for the dissemination of the results. You will also actively participate in the coordination of other studies carried out by the rhinology section within the otorhinolaryngology service of the Marqués de

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Fundación Instituto de Investigación Marqués de Valdecilla CIF: G 39788773

CSV : GEN-245c-5555-dbe8-4b77-30cc-ba8d-8f6e-347f DIRECCIÓN DE VALIDACIÓN : https://portafirmas.redsara.es/pf/valida FIRMANTE(1) : FRANCISCO GALO PERALTA FERNANDEZ | FECHA : 17/10/2024 11:45 | Sin acción específica









Valdecilla University Hospital, collaborating with the research team in the design and execution of scientific projects.

Her activities will include:

- Management of legal documentation, agreements, contracts, etc.
- Administrative management.
- Monitoring the progress of the project, liaising with researchers from different centres.
- Advising researchers on study monitoring.
- Preparation of the necessary monitoring documentation (technical reports, budgets, etc.).
- Preparation of documentation for justification and, where appropriate, auditing.
- Dissemination of the most important aspects of the project.
- Support in all tasks related to the coordination of the CRAIS project.

| PRINCIPAL INVESTIGATOR / RESPONSABLE | RESEARCH GROUP | RESEARCH PROJECT |
|---|--|--|
| Jaime Viera Artiles | INVESTIGACIÓN E INNOVACIÓN EN CIRUGÍA /ORL Rinología | SANOFI23/29: Clinical-radiological algorithm for optimising treatment in patients with chronic eosinophilic rhinosinusitis with poorly controlled nasosinusal polyposis after surgery. |
| | RECRUITMENT INFO | RMATION |
| | | |

| SELECTION PROCESS STAGES (2) | EMPLOYMENT EXCHANGE |
|--|------------------------|
| Admission of applications. Competition phase. Interview pase: maximum number of candidates to be interviewed: 4. Minimum score for this phase: 50. Report of the Tribunal. Resolution. | NOT |
| <u>Note:</u> in order for candidates to be considered for recruitment and employment exchange purposes, they must have a total score of at least 30 points. | |
| SELECTION BOARD | |
| | |

- President: Jaime Viera Artiles, Principal Investigator
- Member: Francisco Galo Peralta, IDIVAL Management Director.
- Member and secretary: Maria José Marín Vidalled, Coordinator of IDIVAL's Technological Services.

| VALUATION OF MERITS | | | | |
|---|------------------------|-------------------|--------|---------|
| MERITS | EVALUATION | SCORE | | MAXIMUM |
| Degree in Social Sciences, Degree in Biochemistry, Degree in Pharmacy, Degree in Biotechnology, Degree in Information Sciences (Documentation), Degree in Statistics, Degree in Physics or Mathematics or Degree in Biomedical Engineering. | Supporting document | Merit fullfilment | YES/NO | 5 |
| Master's degree in biomedical research, public health or epidemiology. | Supporting document | Merit fullfilment | YES/NO | 5 |
| Specific training in database management and statistical analysis: Courses or certificates in tools such as SPSS, R, STATA or management of scientific databases | Curricular | Merit fullfilment | YES/NO | 5 |
| Knowledge of research methodology: Studies or complementary training in clinical study design and scientific 5writing. | Curricular | Merit fullfilment | YES/NO | 10 |
| Training in project coordination or clinical | Supporting document | Merit fullfilment | YES/NO | 5 |

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| | MAXIMUM TOTAL SCORE | | | 100 |
|--|------------------------|-------------------|--|----------|
| MAXIMUM TOTAL SCORE IN INTERVIEW | | | | 60 40 |
| MAXIMUM TOTAL SCORE BY MERITS | | | | |
| | F | INAL SCORE | | |
| Good Clinical Practice (GCP) accreditation. | Supporting document | Merit fullfilment | YES/NO | 10 |
| English | Supporting document | Level | -B2: 5 points -C1 or more: 10 points | 10 |
| Experience in research management | Curriclar | Merit fullfilment | YES/NO | 5 |
| Experience in processing administrative documentation for ethics committees and clinical research in general. | Curricular | Merit fullfilment | YES/NO | 5 |
| studies: Courses or certificates on management and coordination of multicentre projects, preferably in the health area. Experience in clinical data collection and processing, database management, coordination of multicentre studies, bibliographic research, writing scientific articles, collaboration in research projects, use of statistical analysis tools | Curricular | Merit fullfilment | YES/NO | 5 |

(1) Not subsanable

(2) See duration of each phase in the document "Selection Process"

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Santander as of the date of electronic signature

Fdo. Francisco Galo Peralta Fernandez

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