







REFERENCE		OPENING DATE	DEADLINE			
PT20/00084		09/09/2024	18/09/2024			
	PROF	ILE REQUIREMENTS				
	EXCLU	SIVE REQUIREMENTS: (1)				
ACADEMIC DEGREE	Degree + oficial application).	master/equivalente (Justificatio	n must be provided with the			
	VAL	UED MERITS /SKYLLS				
FURTHER	-Training in Phar -Certification in (-Specialisation in Clinical Pharmacology -Training in Pharmacovigilance -Certification in Good Clinical Practice				
	-Experience as D	ne monitoring of Clinical Trials. Data Manager or Study Coordina	tor in Clinical Trials.			
LANGUAGES	-English -Availability for e	external monitoring				
	· · ·	RACT INFORMATION				
TYPE OF CONTR		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	Full time. 1575 hours per year (aprox. 35h/week)			
ANNUAL GROSS S	ALARY	DURATION	OF THE CONTRACT			
2.351, 43 € without prejudice pdate established in state legi			luration of the project or to external n public grants in full competition).			
WORK LOCATI	ONS	UNIT,	UNIT/DEPARTMENT			
Marqués de Valdecilla Univ Pavilion 15-2		Área de Ensayos Clínicos	. Servicio de Farmacología Clínica			
		JOB DETAILS				
	0	FFER DESCRIPTION				
	Rese	arch support technician				
		FUNCTIONS				
he project manager. Ensure effective communic o carry out the start-up vis To carry out the monitoring Verify that the protocol and Ensure compliance with Ge rocedures. Performing the clinical rese Prepare the Monitoring and Keep the essential project of To carry out external monit Support for the resolution queries). Ensure traceability of medio Maintain the information re	ation between the it and train the re in compliance w lits modifications ood Clinical Prac earch project clos Review Reports documentation up coring in centres p of inconsistencient cation delivered t	e research team/promoter in esearch team in the activities with the Monitoring Plan and r s are complied with. tice standards, applicable le ure visit. to the project manager/prom odated and correctly filed in t participating in the clinical tr es, deviations and errors in to assigned sites. assigned sites in the project	nanual, gislation and Standard Operating noter. the centres assigned to him/her. ials of the platform. the data collected from the trial			

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IY

 Produce periodic safety rep Follow-up on any safety iss 	ues that may occu	r with investiga	itional me	dicinal products or h	neurcal devices	
PRINCIPAL INVESTIGATOR RESPONSABLE	R / RESEAR	CH GROUP	RESEARCH PROJECT			
María del Mar García Sáiz		Clinical Pharmacology Science and Innovation) and co-finance				
	RECRUIT	MENT INFOR	MATION	4		
SELECTION PROCESS STAGE	S (2)				EMPLOYMEN EXCHANGE	
 Competition phase. Interview pase: maximum phase: 10. Report of the Tribunal. Resolution. Note: in order for candidates purposes, they must have a final statement. 	to be considered	for recruitment			NOT	
President: María del Mar Gar Member: Francisco Galo Pera Member and secretary: Mar	alta, IDIVAL Manage	ment Director.	TDIVAL'S T	Fechnological Services		
	alta, IDIVAL Manager ria José Marín Vidalle	ment Director.		Fechnological Services.		
• Member: Francisco Galo Pera	alta, IDIVAL Manager ria José Marín Vidalle	ment Director. ed, Coordinator o		-	MAXIMUM	
Member: Francisco Galo Pera Member and secretary: Ma MERITS Specialisation in Clinical	alta, IDIVAL Managei ria José Marín Vidalle VALI	ment Director. ed, Coordinator o	ITS SCOI	-		
Member: Francisco Galo Pera Member and secretary: Mar MERITS Specialisation in Clinical Pharmacology	alta, IDIVAL Managel ria José Marín Vidalle VALI EVALUATION	ment Director. ed, Coordinator of JATION OF MER	EITS SCOI ilment	RE	MAXIMUM	
Member: Francisco Galo Pera Member and secretary: Ma MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical	alta, IDIVAL Managel ria José Marín Vidalle VALI EVALUATION Accreditation	ment Director. ed, Coordinator of JATION OF MER Merit fullf	ITS SCOI ilment ilment	RE YES/NO	MAXIMUM 20	
Member: Francisco Galo Pera Member and secretary: Mai MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical Practice Experience in the monitoring	alta, IDIVAL Managel ria José Marín Vidalle VALU EVALUATION Accreditation Curricular	ment Director. ed, Coordinator of JATION OF MER Merit fullf Merit fullf	ITS SCOI ilment ilment ilment	RE YES/NO YES/NO	MAXIMUM 20 10	
Member: Francisco Galo Pera Member and secretary: Ma MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical Practice Experience in the monitoring of Clinical Trials. Experience as Data Manager or Study Coordinator in	alta, IDIVAL Manager ria José Marín Vidalle VALI EVALUATION Accreditation Curricular Accreditation	ment Director. ed, Coordinator of JATION OF MER Merit fullf Merit fullf	ITS SCOI ilment ilment ilment ilment	RE YES/NO YES/NO YES/NO	MAXIMUM 20 10 5	
Member: Francisco Galo Pera Member and secretary: Ma MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical Practice Experience in the monitoring of Clinical Trials. Experience as Data Manager or Study Coordinator in Clinical Trials.	alta, IDIVAL Manager ria José Marín Vidalle VALI EVALUATION Accreditation Curricular Accreditation Curricular	ment Director. ed, Coordinator of JATION OF MER Merit fullf Merit fullf Merit fullf	ITS SCOI ilment ilment ilment ilment	RE YES/NO YES/NO YES/NO YES/NO	MAXIMUM 20 10 5 10	
Member: Francisco Galo Pera Member and secretary: Mai MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical Practice Experience in the monitoring of Clinical Trials. Experience as Data Manager or Study Coordinator in Clinical Trials. English Availability for external	alta, IDIVAL Manager ria José Marín Vidalle VALI EVALUATION Accreditation Curricular Accreditation Curricular Curricular	ment Director. ed, Coordinator of JATION OF MER Merit fullf Merit fullf Merit fullf Merit fullf Merit fullf	ITS SCOI ilment ilment ilment ilment	RE YES/NO YES/NO YES/NO YES/NO YES/NO -B1: 2 points -B2 or more: 5	MAXIMUM 20 10 5 10	
Member: Francisco Galo Pera Member and secretary: Mai MERITS Specialisation in Clinical Pharmacology Training in Pharmacovigilance Certification in Good Clinical Practice Experience in the monitoring of Clinical Trials. Experience as Data Manager or Study Coordinator in Clinical Trials. English Availability for external	alta, IDIVAL Manager ria José Marín Vidalle VALI EVALUATION Accreditation Curricular Accreditation Curricular Curricular Accreditation	ment Director. ed, Coordinator of JATION OF MER Merit fullf Merit fullf Merit fullf Merit fullf Merit fullf	ITS SCOI ilment ilment ilment ilment	RE YES/NO YES/NO YES/NO YES/NO YES/NO YES/NO -B1: 2 points -B2 or more: 5 points	MAXIMUM 20 10 5 10 5 5 5 5	
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(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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