

(An Autonomous Institute of the Department of Biotechnology, Govt. of India)

NCR Biotech Science Cluster, 3rd Milestone, Faridabad – Gurugram Expressway, P.O. Box No. 04, Faridabad - 121001

Rolling Recruitment notice no.: THS/RN/03/2022

- 1. Translational Health Science and Technology Institute (THSTI) is an autonomous Institute of the Department of Biotechnology, Ministry of Science and Technology, Govt. of India. The institute is an integral part of the interdisciplinary NCR Biotech Science Cluster located at Faridabad, and is designed as a dynamic, interactive organization with the mission to conduct innovative translational research and to develop research collaborations across disciplines and professions to translate concepts into products to improve human health.
- 2. THSTI has built several inter-institutional collaborations and connectivity with industry supported by well-trained teams of research and laboratory staff. This foundation has helped pursuit of thematic research programmes which can be broadly categorized as, (a) Infectious diseases and Immunology (b) Maternal and Child Health, (c) Non-communicable disease (d) Multidisciplinary clinical and translational research. These will be strengthened by four core facilities viz. Small Animal Facility, Data Management Centre, Biorepository and Bioassay Laboratory that will serve not only the research programmes of THSTI, but also the National Capital Region Biotech Science Cluster and other academic and industrial partners.
- 3. This recruitment is to fill up the vacancies under Clinical Development Services Agency (CDSA) of THSTI. CDSA is a niche centre of THSTI established to facilitate development of affordable healthcare products for public health diseases. It is the only public Centre in the country created with a mandate to support and nurture cost-effective, high quality, not-for-profit technology-based preclinical and clinical product development as well as support clinical research conducted by public agencies. It works towards development of an eco-system for training and learning and work with public sector institutions, and small and medium enterprises (SME) to translate innovative technologies into medical products for public good.

The main objectives of CDSA are

- As an academic Clinical Research Unit, to undertake & provide end -to- end clinical study support for investigators and SMEs in study planning, set up, conduct: project management, monitoring, data management, safety reporting, analysis and report writing
- b) Build research capacity and capability through high quality training in the area of clinical development/trials and regulation
- c) Support and strengthen clinical research environment in the country
- d) Regulatory science and policy support: provide tools and approaches to support researchers, regulators, health policy makers & industry
- 4. Applications are invited from eligible candidates to fill up the following positions:

Dated: 11th January 2022

1	Name of the post	Head Regulatory Science and Medical Affairs
	Number of post	One Post
	Emoluments	Up to Rs 1,65,000/-
	Age	55 years
	Minimum Educational Qualification and Experience	Medical professional qualification (MD OR MBBS or equivalent qualification) from a recognized university with at least 10 years of work experience in clinical research especially in clinical operations (start- up activities), regulatory function, medical affairs including medical monitoring, medical writing, pharmacovigilance and medical coding and systems for adverse event review and reporting, safety reporting and management.
	Job profile	Lead the medical and regulatory aspects of clinical trial/studies. Overall responsibility to lead the team on development of protocol, study design, regulatory pathway, medical affairs and safety reporting.
		Serve as medical liaison to all stakeholders – funding agencies, investigators, project teams.
		Leadership and Strategy:
		 Provide leadership on medical and regulatory aspects of clinical trials and clinical study projects. Participate in business development activities Contribute to drafting policies and standard operating procedures Contribute to developing the clinical trials / studies portfolio Act as a key advisor on regulatory matters, and writing position papers as well as advising on "higher risk" studies. Responsible for dissemination of information for CDSA staff on all CDSCO/ ICMR Clinical Trials Regulation and its implications, regulatory requitrements, research governance and Good Clinical Practice (GCP) Serve as a liaison for medical and regulatory functions to all stakeholders – funding agencies, sponsor, investigators, project teams and provides medical and regulatory guidance throughout the life cycle of trials/studies. Provide guidance and oversees safety management, medical monitoring/coding and medical writing functions. Responsible for start-up activities inclusive of regulatory submission dossiers, wherever applicable, and managing the regulatory compliance of the clinical studies. With the Administrative Manager and Chief of Clinical Portfolio Management, oversee and draft Memorandum's of Understanding (MoU's) or other documents to outline the delegation of duties from the sponsor office to CDSA and other stakeholders of the projects Medical Affairs (50%)

- Provide leadership to clinical trial and clinical study projects on medical and safety aspects.
- Participate in clinical review meetings and document preparation for meetings as required
- Guide the project teams in the preparation and review of study documents like oclinical protocols, informed consent forms etc.
- ointegrated clinical and statistical summary reports,
- omeeting presentations
- otherapeutic area training material
- ojournal articles, and other documents
- Review all documents assigned for scientific/ medically relevant issues including drug safety
- Review and sign off technical documents written with respect to medically relevant matters with particular attention to those relating to drug safety
- DSMB: Develop/ review DSMB charter, support constitution of DSMB for clinical trials, organize and coordinate DSMB meetings
- Provide input as necessary to Feasibility Studies, Data and Safety Monitoring Committees (DSMC) and other committees, clinical/ product development planning meetings
- Act as medical liaison with clients and solicit expert advice, develop collaborative relationship with key experts and investigators
- Assist in the preparation of client proposals
- Oversee the medical monitoring and medical coding function for all the clinical studies in which CDSA is involved.
- Train/mentor and provide leadership to the medical monitor(s) and coders assigned to the clinical studies/trials
- Oversee and ensure accurate interpretation of single and or grouped adverse events, serious adverse events, drug effect and attribution of causality, and disease condition
- Oversight / review of clinical narrative reports prepared by the Investigators describing the event; advise on individual participant cases as identified by the study team and identifying queries for the local monitors to complete.
- Ensure compliance with clinical safety and good pharmacovigilance practices and requirements
- Review and provide support in finalising Periodic Safety Update Reports (PSURs)
- Review and edit CSR for clinical consistency with data and standard of practice
- Review and sign off Data Management listings of safety data (including adverse events, laboratory data, vital signs data, medical history, physical examination, concomitant medication),
- Assist the PI and DSM in establishing the presence or absence of clinically meaningful trends and, if noted, assisting in follow up as appropriate with the project team, sponsor, and Regulatory Authorities
- Review and provide input for AEs (coded), past medical history, concomitant medications or other medical data listings to verify and medically vet clinical data.
- Provide consulting services to assist in the development of new drugs or devices under the direction of the Head of Organisation

		Regulatory Science (40%)	
		 Act as regulatory lead to projects by coordinating regulatory work flow for DCGI and Institutional Ethics Committee submission and approvals, ensuring sufficient regulatory project coverage, providing regulatory support to the team. Review and approve investigator site regulatory package documents (Statement of Investigator, investigator CVs, IRB/IEC approval documentation, consent forms, etc.). Work with the appropriate project team members to resolve queries. Maintain a working knowledge of, and assure compliance with, applicable ICH & CDSCO Guidelines, Regulatory Agency requirements, and CDSA SOPs. 	
2	Name of the post	Lead- Clinical Science	
	Number of post	One post	
	Emoluments	Up to Rs 1,20,000/-	
	Age	45 years	
	Minimum	MD from a recognized university with at least 3 years of work experience.	
	Educational	OR	
	Qualification and Experience	MBBS OR BDS or MPH from a recognized university with at least 6 years of work experience in clinical research especially in clinical operations, study planning and	
	Ехрепенсе	conduct, regulatory function, medical affairs including medical monitoring, medical writing, pharmacovigilance and medical coding and systems for adverse event review and reporting, safety reporting and management.	
	Job profile	This position is responsible for taking lead during planning stage of assigned clinical studies and trials, development of grant application in terms of study design, study population, safety reporting, budgeting; protocol development. Timely delivery of key tasks, while maintaining high quality standards are key responsibility areas. The lead Clinical Science will manage the performance of the project team(s) working on projects under his/her direction. Mentoring and development of the project team is a key outcome area for this role. The lead Clinical Science will serve as a point of contact for the sponsors and build sponsor relationships. The lead Clinical Science is also responsible for working cross functionally and understanding the implications of management of clinical science and regulatory affairs activities on other groups within the organization. In addition, this role may also have responsibility for managing the regulatory and medical affairs aspect in clinical/ non- regulatory trial directly. The lead Clinical Science will have direct line reports The post will work closely with the Director of CDSA and Head of Medical Affairs and Regulatory Science.	

3	Name of the post	Biostatistician
	Number of post	One post
	Emoluments	Up to Rs 1,10,000/-
	Age	45 years
	Minimum Educational Qualification and Experience	 MSc in Statistics Minimum 3 years' experience in handling/analysing clinical trial & epidemiological study data Experience in using statistical software, especially base, statistics, and graphics; developing statistical analysis plans and statistical methods for protocols; demonstrated use of sample size estimation and power calculation tools/software Knowledge of clinical data management system (paper based and electronic data capturing)
	Job profile	Responsible for providing statistical support (design, analysis, and reporting) for large scale, clinical trials/studies and/or explore methodological improvements in trial design and conduct. The role includes generating randomization schedules, participating in protocol and Statistical Analysis Plan (SAP) development, preparing and reviewing shell tables, listings, and figures, and reviewing case report forms (CRFs), and database structures.
4 Name of the post Lead- Data Science		Lead- Data Science
	Number of post	One Post
	Emoluments	Up to Rs 1,10,000/-
	Age	45 years
	Minimum Educational Qualification and Experience	Post graduate degree from a recognized university preferably in Computer application/ Computer science/ Data science or relevant branch with experience of working in a research or healthcare environment and atleast 6 years of work experience in clinical data science and/ or data management OR
		Post graduate professional degree preferably in Computer application/ Computer science/ Data science or relevant branch with experience of working in a research or healthcare environment and at least 5 years of work experience in clinical data science and/ or data management

OR

Graduate degree preferably in Computer science/ Computer application or relevant branch with experience of working in a research or healthcare environment and at least 8 years of work experience in clinical data science and/or data management

OR

Professional graduate degree preferably in Computer science/ Computer application or relevant branch with experience of working in a research or healthcare environment and atleast 7 years of work experience in clinical data science and/ or data management

- Working knowledge in software design and development, techniques, testing and validation methodologies and software documentation.
- Strong IT skills
- Experience in Clinical Data Management, Database Administration and Software Development
- Sound working knowledge of clinical database development and monitoring tools and logics & techniques
- Demonstrated experience in preparation of Clinical Study Data Management documents
- Demonstrated experience in software validation and documentation

Job profile

This position is responsible for taking lead during planning of data management for assigned clinical studies and trials, development of grant application in terms of data management, data protection and data security; budgeting for data management. Timely delivery of key tasks, while maintaining high quality standards are key responsibility areas. The Data Scientist will manage the performance of the data management team(s) working on projects under his/her direction. Mentoring and development of the data management team is a key outcome area for this role. The Data Scientist will serve as a point of contact for the sponsors and build sponsor relationships. The Data Scientist is also responsible for working cross functionally and understanding the implications of data management activities on other groups within the organization. In addition, this role may also have responsibility for data management in clinical/non-regulatory trial directly. The Data Scientist will have direct line reports like, but not limited to data manager, Quality Analyst, data coordinator and data entry operator.

- > Call for applications will remain open till suitable candidates are found.
- ➤ Deadline for receipt of applications for each quarter: January 24th, April 24th, July 24th, October 24th 2022.

GENERAL TERMS & CONDITIONS:

- a) For positions at Sr. No. 1, the incumbent will be permitted to undertake consultancy services on behalf of the institute and retain a percentage of the consultancy fees as per the Bye-laws of the institute.
- b) The positions will be hired initially for two years with a probation period of six months. The extension will be granted subject to satisfactory performance of the incumbents and tenure of the project for which they are selected. Those appointed to these positions will not have any claim for regularization of their employment.
- c) All educational, professional and technical qualification should be from a recognized Board/University.
- d) The experience requirement specified above shall be the experience acquired after obtaining the minimum educational qualifications specified for the post.
- e) Closing date of online application will be the **CRUCIAL DATE** for determining eligibility with regard to age, essential qualification etc.
- f) The age limit, qualification, experience and other requirements may be relaxed at the discretion of the competent authority, in case of candidates who are otherwise suitable. In case candidates are not found suitable for the posts notified, they can be offered lower post / lower emoluments on the recommendation of the Selection Committee.
- g) Age and other relaxations for direct recruits and departmental candidates: 1. By five years for candidates belonging to SC/ST communities. 2. By three years for candidates belonging to OBC communities. 3. For Persons with Benchmark Disabilities (PWBD) falling under the following categories: (i) UR ten years, ii) OBC 13 years (iii) SC/ST 15 4. Age is relaxable for Central Government servants up to five years in accordance with the instructions or orders issued by the Central Government, from time-to-time. 5. There is no upper age limit for the Institute employees who are treated as departmental candidates.
- h) Age relaxation as per government norms will be provided duly ensuring at least 5 years remaining service for superannuation (60 years).
- i) All results will be published on our website and all future communications will be only through email
- j) In case a large number of applications are received, screening will be done to limit the number of candidates to those possessing higher/relevant qualification and experience.
- k) With regard to any provisions not covered in this notification, the bye laws of THSTI / Govt. of India rules / guidelines shall prevail.
- l) Canvassing in any form will be a disqualification

HOW TO APPLY:

- 1. **Documents to be kept handy before filling up the online application:** (all the documents except (i) should be in pdf format):
 - i) A soft copy of your passport size photo and signature. (jpeg/jpg/png format)
 - ii) A comprehensive CV containing details of qualification, positions held, professional experience / distinctions etc.
 - iii) Matriculation certificate (equivalent to 10th Standard) / Mark sheet
 - iv) Intermediate certificate (equivalent to 12th Standard) / Mark sheet
 - v) Graduation/Diploma degree certificate / Mark sheet
 - vi) Post-Graduation degree certificate & Mark sheet (if applicable)
 - vii) PhD/MD Degree (if applicable)
 - viii) Relevant experience certificates (if applicable)
 - ix) Caste / Disability certificate in the format prescribed by the Govt. of India, if applicable

2. Procedure for filling up online application:

- i) The eligible and interested candidates may apply online at the Institute's website www.thsti.res.in/career. Applications through any other mode will not be accepted.
- ii) The following will be the step wise procedure-
 - A) Step 1: Details of applicant
 - B) Step 2: Uploading of documents
 - C) Step 3: Payment of application fee
 - The payment can be made by using Debit Card / Credit Card / Internet Banking / UPI.
 - Once payment is made, no correction / modification is possible
 - Candidates are requested to keep a copy of the provisional receipt forfuture reference.
 - Fee once paid shall not be refunded under any circumstances.
 - > Details of fees to be paid are as shown below:

S. No	Applying on direct recruitment	Application fee amount
1.	Unreserved, OBC & EWS candidates	Rs 590/-
2.	SC/ST/Women/PwBD	Rs 118/-

D) Step 4: Submission of application form

- iii) On successful submission of application, an auto-generated email containing the reference number will be sent to the email address provided. Please keep a note of the reference number for future correspondence.
- iv) Candidates are required to keep a printout of the online application form by using the print button on the dashboard for future reference.
- v) Candidates must ensure that he / she fulfils all the eligibility criteria as stipulated in the advertisement. If it is found that he / she does not fulfil the stipulated criteria during the recruitment process, the candidature of the candidate will be cancelled. If the same is noticed after the appointment, the candidate will be terminated following due process.

- vi) Incomplete applications shall be summarily rejected and no correspondence in this regard shall be entertained.
- vii) In case of difficulty in filling up the online form, please send e-mail to **personnel@thsti.res.in** along with the screenshot of the error displayed (if any).

"Government strives to have a work force which reflects gender balance and women candidates are encouraged to apply"

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