

NOTICE OF VACANCY (Plantilla Position)  
  
Center for Drug Regulation and Research

Number of Vacancy	Item Number	Position Title	Salary Grade	Basic Monthly Salary	Civil Service Commission (CSC) Minimum Qualification Standard
1	OSEC-DOHB-MDSP2-3-2000	Medical Specialist II	23	Php 76,907	<b>Education:</b> Doctor of Medicine <b>Experience:</b> 1 year of relevant experience <b>Training:</b> 4 hours of relevant training <b>Eligibility:</b> RA 1080

End User’s Preference:

- Education:** Doctor of Medicine – a graduate of an accredited medical school who finished residency and training specialization  
\*Preferably Oncologist, Pharmacologist, Specialization in Infectious Disease, Cardiologist, Masters in Public Health, Masters in Statistics or Biostatistics, MS in Clinical Epidemiology
- Experience:** **One (1) year** of relevant experience  
\*Preferably with specialization in field of medicine in an accredited training in the chosen specialty with background in Pharmacology, Pharmacotherapy, Toxicology, Epidemiology, Pharmacovigilance (PV), Adverse Events Following Immunization (AEFI), Causality Assessment, Preclinical and clinical researches related to drug development.
- Training:** **Four (4) hours** relevant training  
Preferably in:
- Pharmacology, Pharmacotherapy, Toxicology, Epidemiology, PV/AEFI Causality Assessment, Protocol Development and Evaluation, Clinical Research, Good Clinical Practice
  - Undergone postgraduate training program in a local or International Academic Institution of Merit
  - Member/fellow/diplomate of good standing and certified accordingly by the appropriate specialty society
- Eligibility:** Career Service (CS) Professional/ Second Level Eligibility

Job Description:

(Product Research and Standard Development Division (PRSDD) - Clinical Trial Unit)

1. Review and recommend for approval of clinical trials including amendments, pre-clinical and clinical study reports for new drug application, rationale of fixed-dose combination, product classification and re-classification (e.g. Rx to OTC, initial, MR), Import Permit (Clinical Trials and One- time Importation), Compassionate Special Permit and New Dosage

- Form/ Strength/ Presentation/ etc. of pharmaceutical products. Conduct researches or literature review and/ or peer review on drug product safety and efficacy.
2. Review and recommend GCP inspection findings and report.
  3. Develop policies, guidelines and operational procedures related to the product registration and/ or conduct of clinical trials under the Center's jurisdiction.
  4. Initiate development of systems and approaches on product registration and evaluation of clinical studies including interpretation of results.
  5. Provide technical assistance related to drug product registration and conduct of clinical studies, with interpretation of data or results to stakeholder.
  6. Establish inter/ intra collaboration in the registration of new drug products and in approval and monitoring of clinical studies for FDA purposes.
  7. Provide technical supervision to staff.
  8. Perform other related functions as may be assigned

All qualified next-in-rank personnel shall be automatically considered candidates for promotion. For all interested FDA regular employees including the qualified next-in-rank candidates, they are required to submit item numbers 1, 2, 3, 5, 6, and 7 listed hereunder, to the FDA-Human Resource Development Division (HRDD). Failure to do so shall be deemed a waiver of their right to be included as candidates for the positions applied for. Further, please be advised that those employees with at least Very Satisfactory (VS) rating in the last two (2) Performance Rating periods shall be considered for promotion.

Other interested applicants shall submit the following documents to Food and Drug Administration – Human Resource Development Division for initial documentary review and evaluation:

1. Application Letter with the specific position applied for (Please indicate Item Number);
2. Two (2) sets of duly accomplished Personal Data Sheet (CS Form 212) (downloadable at [www.csc.gov.ph](http://www.csc.gov.ph));
3. Qualification Profile (Annex 1) in 6 copies (downloadable at [www.fda.gov.ph/about/careers](http://www.fda.gov.ph/about/careers))
4. Civil Service (CS) Eligibility/ Board Rating & valid Professional Regulation Commission (PRC) ID;
5. Diploma in any relevant Master's/ Bachelor's Degree and Transcript of Records;
6. Performance Rating for the last two (2) rating periods (for government employees);
7. Certificates of trainings attended, if any (in case of managerial/ supervisory trainings for the last 5 years); and
8. Latest Appointment and Service Record/Certificate of Employment. (FDA Job Order Personnel need not submit this document.

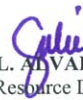

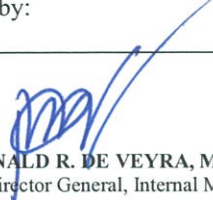
**Note:**

1. Applicants are advised to secure certified true copies/ authenticated copies of documents specified in item numbers 4, 5, and 6 the soonest time, in case of appointment to the position applied for.
2. Applicants are limited to apply up to two (2) vacant positions only.

*\*Online submission of applications are now accepted. ([www.fda.gov.ph/about/careers](http://www.fda.gov.ph/about/careers))*

**Date of Posting:** 17 June 2021

**Deadline of Submission:** 27 June 2021

Prepared by:	Noted by:	Approved by:
 <b>JULIE L. ALVARA, RN, MBA</b> CAO, Human Resource Development Division	 <b>JESUSA JOYCE N. CIRUNAY, RPh</b> Director IV, Center for Drug Regulation and Research	 <b>ATTY. RONALD R. DE VEYRA, MBA, CESO II</b> Deputy Director General, Internal Management