REIMBURSABLE DETAIL Center for Tobacco Products Office of Science

The Center for Tobacco Products (CTP), Office of Science (OS) is offering a Detail opportunity for a **Supervisory Regulatory Health Project Manager, GS-601-14**. The Detail is available immediately for a period up to 120 days. Applicants at the GS-13 and GS-14 level are encouraged to apply. Multiple selections may be made from this announcement. A temporary promotion will be considered.

Area of Consideration:	FDA-Wide
Closing Date:	March 31, 2023
Opening Date:	March 20, 2023
	Beltsville, MD 20705
	11785 Beltsville Drive
	Center for Tobacco Products
	FDA
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Office Location:	Remote (Anywhere in the U.S.)
Position:	Supervisory Regulatory Health Project Manager
Bargaining Unit Status:	Non-Bargaining Unit Position

The CTP, OS offers a fast-paced, dynamic environment and an opportunity to work with dedicated, energetic people who really want to make a difference and improve public health. The position is ideal for someone who wants to have a critical role in the organization and would enjoy the challenge of supervising the success of regulatory health project managers that manage a variety of assignments related to the regulation of tobacco products and CTP workgroups.

Duties include:

The Detail will be located in the OS, Division of Regulatory Project Management (DRPM). Each Branch is responsible for regulatory project management of product application review, research, and program development projects to support implementing the Family Smoking Prevention and Tobacco Control Act. The Detailee serves as a supervisor. Duties may include:

- Supervises staff who coordinate scientific regulatory program coordination, project management, and tobacco application reviews.
- Assigns work to subordinates based on priorities, considering difficulty and requirements of assignments; setting short-term priorities to achieve goals of projects, mentoring, collaborating with team members to identify training needs.
- Monitors and reports actual status of all activities within the assigned projects through interaction with project participants and, if required, supervisors and directors.
- Identifies scientific and regulatory issues, manages them to resolution, and initiates and/or

implements necessary actions consistent with laws and policies.

- Offers suggestions to improve processes, efficiency, and the quality of work of the team.
- Works cooperatively with others, inside and outside the organization, to accomplish objectives, to build and maintain mutually beneficial partnerships, leverage information, and achieve results.
- Serves as the contact point for communications concerning tobacco product applications and other submissions, and ensuring compliance with all legal, regulatory and policy requirements. The employee draws upon a comprehensive knowledge to anticipate and identify subtle or obscure problems and to ensure that the review team is aware of these problems and addresses them.
- Delivers clear, effective communication and takes responsibility for understanding others by asking clarifying questions and summarizing or paraphrasing what others have said to verify understanding.
- Represents the Center in meetings with industry and government agencies during public meetings and venues to provide technical oversight and information on the policies, laws, and regulations that govern the tobacco application review.
- Mentors and trains new and junior-level staff regarding FDA policies and procedures applicable to regulation of tobacco products.

Desired Knowledge and Skills:

- Demonstrate comprehensive knowledge of the Food, Drug, and Cosmetic Act as amended by the Tobacco Smoking Prevention and Tobacco Control Act and related regulations, policies, and procedures.
- Technical knowledge and competency in the areas of basic principles and limitations of biological or physical science, manufacturing, public health policies, and regulations of tobacco products. Skill in applying this knowledge in independently carrying out research or review projects.
- Demonstrated experience effectively leading, planning, and managing projects and resources to accomplish a variety of concurrent activities.
- Prior experience developing policies and programs. Expert knowledge of FDA regulations, statutory authorities, policies, and processes.
- Excellent organizational skills.
- Effective verbal and written communication skills.
- Exceptional interpersonal and team skills as demonstrated by the ability to mentor and train staff, maximize each person's contributions, reconcile divergent viewpoints, and foster collaborative working relationships
- Ability to foster accountability and commitment to the mission of the Center.

Application Procedure:

This Detail opportunity is open to all qualified current FDA employees at the GS-13 and GS-14 grade levels or Commissioned Corps Officers. You must have one year of experience at the GS-13 to be eligible for a temporary promotion. A temporary promotion may be available.

Interested applicants should submit a copy of their resume, most recent copy of SF-50, and statement of interest via email to: <u>CTP-Recruitment@fda.hhs.gov</u>

For questions, please contact Rachel Bartlebaugh <u>Rachel.Bartlebaugh@fda.hhs.gov</u> Please submit a copy of your transcripts, if you are in a different series. Detail is reimbursable. Travel Expenses will not be paid.

Please indicate in the subject line of the email: Detail: CTP, OS, DRPM - Supervisory RHPM, GS-0601-14 – March 2023

If you are not currently in the GS-0601 series, please submit a copy of your unofficial transcripts or a previous SF-50 showing the GS-0601 series.

Detail is reimbursable. Travel Expenses will not be paid.

Candidates must express interest by March 31, 2023.

Supervisory concurrence is required in order to accept a Detail; it is NOT required to apply.

*This is not an official vacancy announcement under the Merit Promotion System.