Company: Pfizer Inc.
Salary range:
Job Title: Director, Clinical Monitor, Vaccines
Brief description of the position: Pfizer Vaccines has a very active vaccine pipeline and is currently developing several vaccines. This position is to join the COVID vaccine program and/or the RSV vaccine program. Both are conducting large vaccine efficacy studies. This position would provide an opportunity to join a very dynamic team as physician/medical monitor and work on clinical trials and other related vaccine development work.

Pfizer’s Pearl River site has been involved in vaccine development for over 100 years. Currently, hundreds of staff at the site support Pfizer’s large vaccine pipeline as well as its approved vaccines creating a large vibrant professional community committed to vaccine development. The site is in commuting distance of Westchester County NY, upper Manhattan, and Bergen County NJ.

ROLE SUMMARY

The clinician medical monitor is accountable for providing medical and scientific expertise and oversight for Clinical Trials and serves as a single point of accountability for design, execution, monitoring, delivery and reporting of one or more clinical studies and to ensure patient safety.
The clinician medical monitor may be required to design a development strategy for multiple protocols designed to obtain worldwide approval for a compound or group of compounds.
In addition to study level activities, the clinician medical monitor may participate in standing committees, review compounds for potential in-licensing, including performance of due diligence reviews, and provide assistance to new business development on market opportunities and the target product profile.

Please visit to see the complete job
Qualifications you are seeking: BASIC QUALIFICATIONS

Candidate demonstrates a breadth of diverse leadership experiences and capabilities including: the ability to influence and collaborate with peers, develop and coach others, oversee and guide the work of other colleagues to achieve meaningful outcomes and create business impact.

Licensed by a health authority to prescribe medicines (independent of supervision) for at least one year (post "intern/houseman" year), and has utilized the license to prescribe medicines in a patient care setting for an aggregate duration of at least one year.
Possesses the ability to critically evaluate medical/scientific information.
Excellent written an oral communication.
Understands the design, development, and execution of clinical programs and studies.
Capacity to adapt to a fast pace and changing environment.
Documented experience in the pharmaceutical industry related to clinical research programs and registration activities.
Responsible for managing multiple studies

PREFERRED QUALIFICATIONS

Medical degree (M.D./D.O. or equivalent).
Documented work experience/knowledge of statistics.
Training and experience in infectious diseases and/or infection control in the hospital setting is preferred.
Experience with investigational clinical trials is preferred

Phone:
Fax:
Address:

Website: