

## QUALITY CONTROL DOCUMENT SPECIALIST

This position will be responsible for activities involving quality assurance and compliance with applicable regulatory requirements within the Quality Control Group.

**ESSENTIAL DUTIES AND RESPONSIBILITIES** include the following. Other duties may be assigned.

- Performs the compliance review and approval of all documentation related to disposition of clinical products and research testing documentation
- Generation of associated deviations, investigations and OOS reports
- Generation of change controls
- Performing routine internal audits of quality control processes
- Scheduling, initiation and review of stability study data
- Assisting in all other QC related activities as required
- Follows Good Manufacturing Practices (GMPs), Standard Operating Procedures (SOPs)
- Other duties as assigned or determined by Area Management

### **EDUCATION and/or EXPERIENCE**

- Must have a Bachelor's degree in life sciences or an Associate's degree with 3-5 years of relevant experience and/or training; or equivalent combination of education and experience
- Experience with in a pharmaceutical/biologics environment is preferred
- GMP experience is a plus

VGXI is an Equal Opportunity Employer

Contract to hire position