



GROUP
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JOB DESCRIPTION

ROLE: Executive - DQA

SALARY AND BENEFITS: Negotiable

REPORTS TO: Asst. Manager - DQA

LOCATION: Vadodara

PRIMARY RESPONSIBILITIES:

- Create and Review of technical documents including analytical method development protocols and reports, standard operating procedures in compliance with regulatory requirements.
- Preparation and Approval of Standard Operating Procedures and ensure the preparation, Issuance, Revision/Review and Retrieval of the documents.

SECONDARY RESPONSIBILITIES:

- Preparation and Approval of Standard Operating Procedures.
- Maintain documentation systems and filing of all relevant records, reports and other documentation associated with projects.
- Preparation of Process Validation Protocols and summary reports based on the Analytical results and Batch documents data.
- To co-ordinate the Qualification / Validation of the Equipment, Systems and Process.
- To ensure the issuance of required documents for Drug and Regulatory Affairs.
- To ensure the preparation, Issuance, Revision/Review and Retrieval of the documents.
- To maintain laboratories to GMP standards.
- Any other duties as required.

ESSENTIAL EXPERIENCE, SKILLS AND ABILITIES:

- Essential Masters in Pharmacy or Masters in Science.
- Required experience for Executive 2-4years.
- Knowledge of Quality Assurance or Analytical Research and Development.
- Knowledge of cGMP/cGLP.
- Experience of working in an QA, AR&D, IPQA or Regulatory audit compliance environment.
- Good knowledge of Validation Documents Review.
- Ability to work on own initiative and in a team.
- Excellent writing and communication skills are must.
- Ability to adapt to change.