



Title: **QC Analyst**

Reporting to: **QC Manager**

### **Role and Responsibilities**

Duties will include but are not limited to the following:

- Responsible for performance of stability testing (real time and accelerated) on Technopath products.
- Work with other team members to develop procedures, test protocols and appropriate acceptance criteria for stability studies.
- Organise materials and monitor stock levels of materials used for stability test program.
- Perform laboratory analysis using various clinical laboratory instrument platforms.
- Perform value assignment testing for Technopath control materials – work with other team members to develop protocols, perform testing and collate data.
- Perform troubleshooting and routine maintenance of lab instruments and equipment
- Document and analyse data, ensuring that all data produced is accurately assessed.
- Ensure work is completed to schedule, as dictated by project plans and production schedules.
- Assist in technical failure investigations as required.
- Carry out testing of raw material, in process or finished products as required.
- Identify areas for continuous improvement

### **Job Requirements:**

- Minimum of diploma qualification in a relevant Science discipline
- 3+ years' experience working in a clinical laboratory or other regulated medical product environment.
- Experience in routine use of clinical laboratory instrumentation.
- Experience in troubleshooting and maintenance of clinical laboratory instrumentation
- Knowledge of GLP requirements

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