

## Director of Quality Assurance – In Vitro Diagnostics

MBio Diagnostics of Boulder, CO ([www.mbiidx.com](http://www.mbiidx.com)) is seeking a Director of Quality Assurance (QA), or Director of Regulatory Affairs / Quality Assurance (RA/QA). MBio is technology-driven company focused on multiplex point-of-care products with medical, food safety, and environmental applications. Our new Quality professional will manage MBio's Quality System and lead the ISO 13485:2016 certification program.

### Job Duties:

- Responsible for the QA activities linked to the company's FDA QSR-compliant design, development, and manufacturing operations.
- Streamline MBio's Quality Management System procedures.
- Lead the company through ISO 13485:2016 recertification with a notified body.
- Draft Standard Operating Procedures (SOPs), educate staff, and perform many QA department functions as the company grows. This will be a "hands on" position.

Demonstrated background in guiding teams through current Good Manufacturing Practice (GMP) development is required. The ideal candidate will have demonstrated QA background in organizations developing point-of-care diagnostics and CLIA-waived devices.

### Expectations and skills:

- Work with MBio Management and Quality consultants to review, revise, and streamline MBio's Quality Management System (QMS) standard operating procedures.
- Work closely and effectively with MBio R&D and Manufacturing teams to create an effective and compliant QMS for an early stage (< 50 employees) IVD product developer.
- Lead the Company's notified body audit preparation and certification process.
- Ideal candidate will have a background in FDA *in vitro* diagnostics product development, and demonstrated skills working with and guiding development teams under Design Control.
- Candidate will have outstanding communications skills and will be able to facilitate discussions between Senior Management, R&D, Engineering, and Manufacturing.
- Candidate will evaluate, recommend, and implement quality management software packages for document control, training records, and potentially other QMS functions.
- ISO 13485 auditor or lead auditor certifications will be considered favorably.

No single candidate is expected to have demonstrated skill in all functions listed below, but candidates should highlight relative experience in a brief cover letter.

- Prior experience developing Quality Management Systems from the ground up, or

significant revisions of systems.

- Prior experience in a small company setting, and related experience with “right-sizing” the QMS.
- Experience and skill with managing QA programs that include software development under IEC 62304.
- Experience and skill implementing Change Control processes.
- Management and documentation of training.
- IVD or medical device manufacturing experience.
- CLIA-waived device development background.

**Education and Experience:**

- Bachelor’s or higher degree in life sciences field.
- A minimum of 7 years IVD / medical device industry experience with a minimum of 5 years in quality assurance.

Applicant must have authorization to work in the U.S. Resumes must be accompanied by a cover letter explaining how the applicant meets the job requirements and desired skills. Please email cover letter and resumes to [jobs@mbiodx.com](mailto:jobs@mbiodx.com) with applicant name and the job title listing in the subject line. No phone calls, please.

MBio Diagnostics, Inc. is an Equal Opportunity Employer.