Q.I. Medical, Inc.

Compounding Testing Products for Regulatory Compliance and Quality Assurance

For over 20 years, Q.I. Medical has been a trusted provider of quality assurance products to compounding pharmacies throughout the world. Our focus is on developing easy to use products for our customers to help them with their in-house testing needs. Whether it is a low, medium, or high-risk facility, Q.I. Medical can help them to be in compliance with regulatory guidelines (USP <797>, USP <71>, 503a or 503b, etc.). Cost effective disposable products are available for aseptic/media fill, sterility testing, environmental monitoring, gloved fingertip sampling, hazardous drug manipulation testing, vial adaptors, and other growth media and filtration related products. Support equipment includes: incubators, filter integrity testers, vial blocks, and UV lights. All media is challenged with a battery of USP specified organisms, and lot specific Certificate of Analysis (CofA) are available for download. All products are sold through regional stocking distributors in order to provide fast local service and support. Please contact us for more information.



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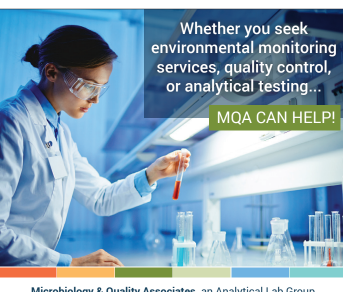
Associates of Cape Cod, Inc.

Your Endotoxin Experts

Specializing in chromogenic and turbidimetric reagent technologies, Associates of Cape Cod, Inc. (ACC) has been a leader in endotoxin detection products and services for more than 40 years. ACC pioneered LAL testing methodology and was the first FDA licensed company to manufacture LAL reagents; ACC has grown to be an internationally recognized leader in endotoxin detection. With a dedication to quality, ACC is certified to I.S. EN ISO 13485:2012 and ISO 13485:2003. We are FDA inspected and operate DEA Licensed and CLIA-certified laboratories. Our endotoxin detection reagents, instruments, and software are used within the pharmaceutical, medical device, biotechnology, compounding pharmacy and dialysis industries for quality control, product release, and research. Our reagents are FDA licensed and can be used for testing in compliance with USP, EP, and JP bacterial endotoxin test chapters and our software is 21 CFR Part 11 Compliant. ACC also operates a Contract Test Services (CTS) Laboratory which has specialized in testing for endotoxin contamination for over 30 years. Our CTS laboratory is GMP compliant, ISO registered and DEA licensed and is capable of handling all controlled drug substances except those included in Schedule 1. All testing services can be performed to FDA, USP, EP, and/or JP regulatory guidelines.



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Microbiology & Quality Associates, an Analytical Lab Group portfolio company, will apply its resources — including a GMP FDA registration, ISO 17025 accreditation and ISO 9001 compliance — to bring clients the best service and most timely, accurate results.



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Quality Science. Quality Service. Quality Results.

Microbiology & Quality Associates Inc., an Analytical Lab Group portfolio company in the San Francisco Bay area, provides contract biotech development services, testing services, consulting, validation and calibration services, training, and scientific resources. MQA is FDA inspected and registered. Our Quality System complies with GxP regulations and we are ISO 17025 accredited and 9001 compliant. MQA staff have many years of experience providing services to compounding pharmacies in the United States. Our philosophy is to do things right the first time, use the best science, and provide top quality results to our clients. We put our clients first by providing outstanding customer service, and flexibility to help you meet your deadlines.



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