QUALITY ASSURANCE MEDICAL DEVICES (QAMD)

QAMD 400

36 Hours

Medical Device Quality Assurance Specialist, Introduction

(Formerly BMGR 645 Introduction to Medical Device Quality Assurance) This course introduces students to the medical device industry in preparation for an entry-level position as a quality assurance specialist. Learn the role and responsibilities of a quality assurance specialist within the medical device industry. Learn about the industry \diamondsuit size and scope, current trends, and products used in healthcare settings. *(Apportionment)*

QAMD 405

36 Hours

36 Hours

Regulatory Affairs for Medical Devices

(Formerly BMGR 648 Regulatory Affairs for Medical Devices) Course will provide students with an in-depth understanding of the regulations and regulatory agencies that are specific to the medical devices industry. The course will cover U.S. and European Union (EU) regulations and related agencies. Includes laws governing the development,

QAMD 410

Document and Database Management for Medical Devices

manufacturing and approval of medical devices. (Apportionment)

(Formerly BMGR 651 Document and Database Management for Medical Devices)

An overview of regulatory requirements for document control and database management for the medical device field. Covers the fundamentals of writing documents that meet regulatory compliance. Equips students for handling document management systems efficiently. *(Apportionment)*

QAMD 415

36 Hours

(Formerly BMGR 654 Quality Assurance for Medical Devices)

This course provides students with an in-depth overview of quality assurance and its role within the medical device industry. The course covers the role of assurance during the manufacturing and production stages. Topics will include the step-by-step process of ensuring quality requirements are met for a product or service. (*Apportionment*)

QAMD 420

36 Hours

36 Hours

Technical Writing for Bio-Medical Industries

(Formerly BMGR 657 Technical Writing for Bio-Medical Industries This course will provide students with an overview and understanding of technical writing for the bio medical industry. The course will cover principle writing methodologies for quality assurance, engineering, manufacturing, and production of a medical device product. Topics will address documentation as related to FDA mandated requirements, ISO standards and for writing GMP procedures. *(Apportionment)*

QAMD 425

Quality Auditing for Medical Devices

Quality Assurance for Medical Devices

(Formerly BMGR 660 Quality Auditing for Medical Devices)

This course presents the principles and techniques for assessing the adequacy of a quality system for a medical device manufacturer. Topics include evaluating the quality system as it conforms to FDA regulatory requirements, standards, review of standard audit terms and other audit concepts. *(Apportionment)*