Registered under The Companies Act, 1956 & Various International Quality Certification Bodies.

Vision Care Certification

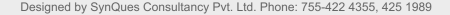
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We Provide global assessment & certification consultants services.

100%



Management Certification Product Certification Inspection **Compliance Auditing** Training Testing

isioncare certification

success rate for ISO certification

ABOUT

Certification Consultancy

Visioncare Certification Pvt. Ltd (VCPL) is a registered company under Registrar of companies 1956 Actvide Registration No.U74200MP2013PTC030645, and also Registered with Trademark Authority, with our head office at Bhopal.Visioncare Certification Pvt. Ltd (VCPL) is a provider of consistent, competent & confidential services for ISO 9001 (QMS), ISMS (Medical Devices), ISO 14001 (EMS), (OHSAS) 18001, ISO 22000 (FSMS), ISO 27000 (ISM), ISO 50000 (Energy) &, GMP, CE Marking, BIFMA, Trade Mark & other branding services.

Values

We, at Visioncare, are committed to achieve total customer satisfaction by providing quality based consulting services that meet and exceed customer expectations. Our focus on continually improving our Quality Management System is our means of achieving operational excellence and customer trust.

Mission

- To provide high quality Consultancy at affordable cost, keeping customer satisfaction as our primary motive
- To provide total Quality solutions at a single door step.
- To promote and support Quality related research at all levels.

SERVICES

ISO 9001:2015 (Quality management Systems)

specifies requirements for a Quality management Systems where an organization needs to demonstrate its ability to consistently provide product that meets customers and applicable statutory and regulatory rewquirements, and aim to enhance customer satisfaction through the effective application of the system, including processes for continual improvement of the system and the assurance of conformity to customer and applicable statutory and regulatory requirements.

ISO 14001:2015 (Environmental Management Systems)

species requirements for an Environmental management Systems to enable an organization to develop and implement a policy and objective which take into account legal requirement and other requirements to which the organization subscribes, and information about signification environmental aspects. It applies to those environmental aspects that the organization identifies as those, which it can control, and those, which it can influence. It does not itself state specific environmental performance criteria.

ISO 27001:2015 (Information Security Management System)

specifies the requirements for establishing implementing, operating, monitoring, reviewing, maintaining and improving documented information. Security Management System within the context of the organization's overall business risks. It specifies requirements for the implementation of information security controls customized to the needs of individual organization or part thereof.

ISO 22000:2005 (Food Safety Management Systems)

is a generic Food Safety Management system Standard. It defines a set of general food safety requirements that apply to all organizations in the food chain. Unlike some standards, ISO 22000 does not follow prescriptive checklist approach. Instead, it allows an organization to develop food safety management Systems that meets the needs of its suppliers and customers. HACCP stands for Hazard Analysis and Critical Control Points. HACCP is an Industry-wide effort approved by the scientific community as well as regulatory and Industry practitioners. It is a Food Safety methodology that relies on the identification of Critical Control Point (CCP's) in food production and preparation process. Closely monitored CCP's will ensure that food is safe for human consumption.

ISO 45001:2018

is an assessment specification for Occupational Health safety obligations is an efficient manner, OHSAS 18001 Certification improves that efficiency of the internal operations, thus reduces the chance, accidents and downtime. The safety and quantity of the employees is the prime responsibility of the organization as the future hazards affect their efficiency. OHSAS 18001 assures the compliance with present legal requisites and reduces the risk of various penalties and possible litigation.

ISO 13485:2016 (Medical Device)

is an ISO standard, published in 2004, that represents the requirements for a Comprehensive management system for the design and manufacture of medical devices. It provides a framework for companies to meet the customer and regulatory requirements. The main goal is to provide a harmonized model for Quality Management System requirement in the International market since different countries might have different standards, MCPL's comprehensive SIO 13485 Audit program for the Medical device Industry enables your organization to implement an effective ISO 13485 Quality management System and meet domestic and International regulations for medical device while achieving overall business improvement.

IATF 16949 : 2016

IATF 16949:2016 (replaces ISO/TS 16949:2009) is a standard that establishes the requirements for a Quality Management System (QMS), specifically for the automotive sector. The ISO/TS 16949 was originally created in 1999 to harmonize different assessment and certification schemes

SA 8000 Social accountability

SA 8000 is based on the Principles of International Human rights norms and described in international Labour organization Conventions, the United Nations Convention on the Rights of the child and the United Nations convention on the rights of the child and universal declaration of human rights. It measures the performance of the companies in eight. It measure the performance of companies in eight key Areas: child labour, Forced labour, Health And Safety, free association and collective bargaining, discrimination, disciplinary practices, working hours and compensation.

CE Marketing

is a symbol of product safety. This is product certification for certain products to be sold in European Union. The letters "CE" are the abbreviation of French Phrase "Conformite Europeene" which literally means "European Conformity". Initially used term was "EC mark" and later on "CE marking" officially replaced it in the directive 93/68/EEC in 1193. "Now CE marking" is used in all EU official documents. If you manufacture or import a product which falls within the scope of one or more of the New Approach directives and wish to place your product on the market in any of the members states of the European Economic Area (EEA), then you must apply CE Marking to your product against the essential requirements of all these applicable directives.

GMP (Good manufacturing Practice)

refers to the Good manufacturing Practice regulations promulgated by the US Food and drug Administration under the authority of the Federal food, Drug and Cosmetic Act. These regulations, which have the force of law require that manufactures, processors and packagers of drugs, medical device, some food, and blood take proactive steps to ensure that their products are safe, pure and effective, GMP regulation require a quality approach to manufacturing, enabling companies to minimize or eliminate instance of contamination, mix-ups, and error. This is turn, protects the consumer from purchasing a product, which is not effective or even dangerous.

Energy Management Systems (ISO 50001)

An Energy management Systems (EnMS) is a systems of computer-aided tools used by operators of electric utility grids to monitor, control, and optimize to performance of the generation and/or transmission systems. The Monitor and control function are known as SCADA; the optimization packages are often referred to as "advance applications"