Vesismin Health’s Quality Policy

The Management of Vesismin, SL, is committed to comply with the Quality Policy and Objectives displayed in this document, as well as the regulatory requirements of Medical Devices (Directive 93/42/EC and Royal Decree 1591/2009) and of Biocide Products (EC Regulation 528/2012, Royal Decrees 3349/1983, 162/1991, 443/1994 and 1054/2002), through the development and improvement of the Quality Management System of Vesismin, SL, according to ISO 9001 and ISO 13485 standards.

The Management is committed to all the people that make up Vesismin, S.L. know and accept the provisions described in the Quality Manual, in order to effectively meet the quality objectives:

- To get the satisfaction of customers and stakeholders.
- To establish a continuous improvement of the Quality System (evaluating said system periodically).

Vesismin, S.L. is dedicated to developing, manufacturing, importing and marketing Medical Devices (disinfectants of invasive medical devices and surfaces of non-invasive medical devices) and disinfectants for clinical and industrial environments. As added value, it provides advisory services and training in accordance with the requirements and needs of our customers. Vesismin complies with applicable legislation and regulations, without accepting any commitment that may affect the quality and efficacy of said products and services, or the safety of patients.

The purpose of Vesismin is to contribute to reduce infections by working honestly and responsibly, taking special care of relationships with stakeholders (employees, customers, the environment, suppliers...) so that they are aligned with our values and purposes.

Vesismin, S.L.

Víctor Vallés
General Manager

July 2018