

DOCUMENTATION
INFORMATION
statistics



sterile good- and
lot documentation





GOAL AND FUNCTIONALITY

KCC Steri-Doc grants a smooth run as well as a complete documentation of individual activities and machine programmes within the sterile good preparation and – management.

The KCC Steri-Doc records the realization including the following facts: WHO (which staff), WHAT (which medical device—at which individual set level), WHEN (at which time), WHERE (at which workstation), WHERE-BY (with which machine, which programme or instrument) and HOW (sufficient, insufficient).

It is identifiable at any time when which sterile good is at which preparation site in the AEMP.

PROGRAMME PARTS

- user management
- master data module
- recording of orders and sterile good documentation in the preparation process
- interface for the import of machine-data
- preparation verification
- module for statistics



consistent sterile
good documentation



INFORMATION AND OVERVIEW

DEVELOPMENT

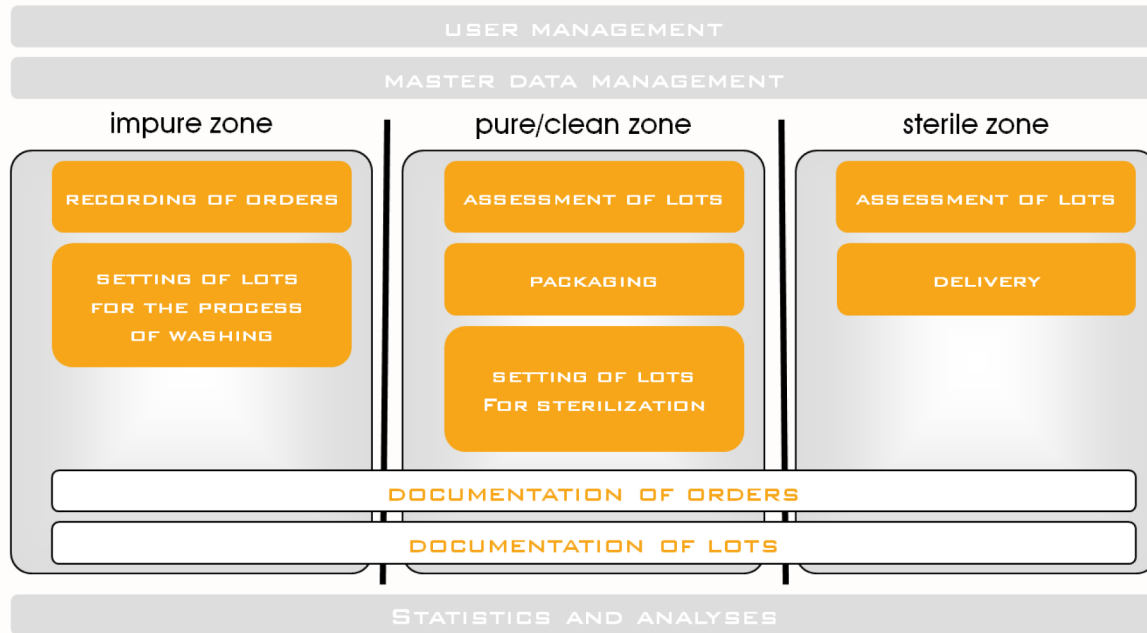
The software is subject to a continuous development and re-implementation. These changes will be added to the existing programme after the development in the course of a maintenance agreement.

An adjustment to further processes of the sterilization is possible due to a very flexible design of the software.

Specifications of the system:

- basis is MS Windows 32 Bit,
- relational SQL-database
- client-server application

VENDOR- AND MACHINE INDEPENDANT



ADVANTAGES

- An extensive user management enables a smooth circulation of personal employed and it also protects programm individual critical settings against unauthorized changes.
- Individual components and processes can be adapted quickly and easily to appropriate needs due to various programme settings.
- There is the possibility to adopt medical devices of excel-sheets and (producer-) catalogues regarding master data as far as they are available in a compatible way.
- The workflow can be executed via barcode scanner almost completely.
- The machine data which are imported via an interface are being diagrammed and recorded.
- The reference to the patient can be made through labeling of the sterile good sets with a special number for unambiguous assignment (which set has been used for which patient).

FLOW VIA
BARCODE SCANNER, RFID





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