



## SERVICES QUALITY MANAGEMENT MEDTECH

**The set-up and maintenance of a quality management system according to ISO 13485 / QSR are an indispensable requirement to be active in the medical engineering sector. We support you in all important steps of a (re)certification or adaptation of the QMS to new requirements and offer you comprehensive services.**

### Your contact

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### Your Advantage

You can benefit from our vast experience as a self-certified and practically consulting company as well as from the practice know-how of some employees as experts/auditors. With our tailor-made service you decide on the pacing and the volume of our services, from a non-recurring coaching through to the extensive outsourcing of the quality management tasks.

### Our Services

#### Coaching

Coaching during the setting up of the QMS to the point of support and review of certification. This includes in particular:

- Identification of the processes to be described
- Support in the interpretation/implementation
- Monitoring/support in the preparation of expertise of system and the certification audits
- Formulation of the conform description of the value-added process with focus on „design and development“ including risk management

#### Design dossiers / Technical documentation of products

Compilation of GAP–analyses related to design dossiers based on the latest CE and FDA requirements

- Active support in closing of possible discrepancies
- Review in terms of content / revision of dossiers

#### Q-Tool® as a possible document management system

Based on our own experience we have developed a document management system „Q-Tool“ which is tailor-made to the requirements of ISO 13485 and enforces discipline in a simple way without restriction of the flexibility you are accustomed to by Office Suite. Please see our separate factsheet about Q-Tool®.