

上海波西认证有限公司

Shanghai POSI Certification Co., Ltd.

ISO 13485 医疗器械质量管理体系认证过程管理程序

**ISO 13485 Medical Device Quality Management
system Certification Process Management Procedure**

文件编号 Doc. No.: POSI/P07

版本 Version: 07

编制 Produced by: 认证部 Certification Dept.

修订 Revised by: 认证部 Certification Dept.

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初始发布:
Initial Issue

2018-05-03

修订日期:
Revision Date

2025-08-06

实施日期:
Implementation Date

2025-08-06

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