朝阳小红门北京医疗器械三类经营许可证二三类专业快捷包拿证

产品名称	朝阳小红门北京医疗器械三类经营许可证二三类专业快捷包拿证
公司名称	北京星期三企业管理咨询有限公司业务部
价格	.00/件
规格参数	医疗器械三类:注册 医疗器械二类:注册 注册公司:网络销售备案
公司地址	北京市海淀区清河嘉园东区甲1号楼11层1124
联系电话	15501182773 15501182773

产品详情

- (2) 相关部门受理申请人的申请;
- (3) 到实际场地进行勘察以及对产品进行审核;
- (4) 准予颁发三类医疗器械许可证。
- 2、法律依据:《医疗器械监督管理条例》第十四条

第一类医疗器械产品备案和申请第二类、第三类医疗器械产品注册,应当提交下列资料:

- (一)产品风险分析资料;
- (二)产品技术要求;
- (三)产品检验报告;
- (四)临床评价资料;
- (五)产品说明书以及标签样稿;
- (六)与产品研制、生产有关的质量管理体系文件;
- (七)证明产品安全、有效所需的其他资料。

法律责任:

申请人以欺骗、贿赂等不正当手段取得《医疗器械经营企业许可证》的,(食品)药品监督管理部

elevant departments shall accept the application of the applicant; (3) Investigate the actual site and audit the products; (4) Grant the issuance of a class III medical device license. 2. Legal basis: Article 14 of the Regulations on the Supervision and Administration of Medical Devices As for the registration of Class II and III medical devices, the following materials shall be submitted: (1) Product risk analysis data; (2) Technical requirements for the products; (3) Product inspection report; (4) Clinical evaluation data; (5) Product specification and label samples; (6) quality management system documents related to product development and production; (7) other materials required to prove the safety and effectiveness of the product. legal liability: Where the applicant obtains the License of Medical Device Trading Enterprise by cheating, the (food) and Drug Supervision and Administration Department