自测版试剂盒CE认证需要准备什么资料

| 产品名称 | 自测版试剂盒CE认证需要准备什么资料 |
|------|-------------------------|
| 公司名称 | 全球法规注册CRO-国瑞IVDEAR |
| 价格 | .00/个 |
| 规格参数 | |
| 公司地址 | 光明区邦凯科技园 |
| 联系电话 | 13929216670 13929216670 |

产品详情

1. 自测版试剂盒CE认证需要准备哪些资料?

答:请参考以下资料清单:

Required documents (Take NB1434 as an example)

| St | The copy of | |
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| at | National Court | |
| е | Register docume | nt |
| | or certificate of | - |
| | entry in the | |
| | register of busine | ~~ |
| | | :00 |
| | activity | |
| | The written | |
| | declaration that r | າວ |
| | application for | |
| С | product | |
| 0 | assessment has | |
| m | been submitted t | 0 |
| ni | another Notified | |
| t | Body | |
| m | The written | |
| е | declaration that r | າວ |
| | medical incident | |
| | involving the | • |
| Ű | medical device | |
| | applied for | |
| | certification have | |
| | | ; |
| | been occurred | |
| | The | |
| | Manufacturer ' a | S |
| | commitment to | |
| | maintain the | |
| | procedures of | |
| | regular review of | |
| | experience gaine | d |
| | with the in | |
| | vitro diagnostic | |
| | medical device | |
| | subsequent to its | |
| | being placed on | |
| | the market | |
| | The | |
| | | 5 |
| | commitment to | 2 |
| | | |
| | fulfil the | |
| | obligations | |
| | resulting from his | |
| | Quality Assurance | ce |
| | System | |
| | | |
| | | |

| | The |
|---------|----------------------|
| | Manufacturer's |
| | commitment to |
| | maintain the |
| | effectiveness of the |
| | Quality Assurance |
| | System |
| | The |
| | Manufacturer's |
| | commitment to |
| | maintain the |
| | procedure in |
| | accordance with |
| | the chapter 9 of |
| | the Act on |
| | Medical Devices |
| | (the Polish Act) in |
| | case of receiving |
| | an information on |
| | medical incident |
| | The Authorised |
| | Representative 's |
| | written statement |
| | about cooperation |
| | with Manufacturer |
| | (if applicable) |
| | The description of |
| | the type of |
| | product including |
| | variants and the |
| | list of differences |
| | between variants |
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evThe essential ic requirements e checklist (Annex 1 to the Regulation of the Minister of Health of 17 February 2016 on essential requirements and conformity assessment procedures of in vitro diagnostic medical devices as amended (Journal of Laws of the Republic of Poland 2016, item 211) The design drawings, the specifications of components and parts, circuit diagrams The results of the design calculations Diagram of the manufacturing process including the midproduction inspection with indication of subcontracted stages The list of harmonized standards applied in whole or in the part and other applied standards Description of meeting the essential requirements, if they are not based only on the harmonized standards

Test reports showing conformity with requirements of harmonized standards and/or Common Technical Specifications Performance evaluation data confirming parameters declared by the manufacturer Stability testing data confirming the stability declared by the manufacturer Report of testing with participation of nonprofessional users (if applicable) Declaration of conformity of the in vitro diagnostic medical device Risk management and risk analysis, including: Report of risk analysis for conformity with PN-EN ISO 14971/EN ISO <u>14971</u> A draft marking (label), package (in case of translation of information documents, a formal confirmation from translation agency shall be attached)

Instructions for use of a medical device (in case of translation of information documents, a formal confirmation from translation agency shall be attached) Information on the in vitro diagnostic medical device restrictions Information on the origin and conditions under which human tissues are derived or substances derived from these tissues - refers to medical devices containing human tissues or substances derived from these tissues Brochures, folders, presentations and other promotional materials concerning in vitro diagnostic medical devices prepared by the Applicant (in case of translation of information documents, a formal confirmation from translation agency shall be attached)

| | List of all suppliers |
|----------|-----------------------|
| | and |
| | subcontractors, |
| | indicating key |
| | 0, |
| | critical suppliers/s |
| | ubcontractors; |
| | certificates of supp |
| | liers/subcontractor |
| | s (if applicable) |
| | Product sample to |
| | be consulted with |
| | the PCBC |
| M | Quality Manual + |
| | Quality Policy and |
| | Objectives |
| | Organizational |
| | chart and |
| | |
| | responsibilities |
| | and competencies |
| | of the |
| | management |
| u | Document and |
| ali | record control |
| ty | procedure |
| A | Risk management |
| | procedure |
| | Designing |
| | procedure |
| | Manufacture |
| | procedure |
| | Procedure for the |
| | |
| 1. | purchasing and |
| | subcontractors ' |
| | control |
| | Non-complying |
| | device control |
| | procedure |
| u | Corrective and |
| m | preventive actions |
| e | procedure |
| | Servicing |
| | procedure (if |
| | applicable) |
| | Sterilization |
| | procedure (if |
| | r ` |
| | applicable) |
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In vitro diagnostic medical device identification and traceability procedure Product security procedure Measuring equipment supervision procedure Procedure for obtaining the feedback from users concerning product (data analysis) Procedure for the testing of in vitro diagnostic medical device during the manufacture process or/and testing of final product Internal audit procedure Product measurement procedure Procedure for issuance and implementation of advisory notes Procedure for dealing in case of medical incidents Procedure for regular review of experience gained with the device subsequent to its being placed on the market