# **≫ 检查手套(北瑞)** 产品编码: LEG-BR

检查手套,无粉麻面,采用进口优质天然胶乳加工而成,适用于低风险的医疗护理、医疗检查、卫生防护等,可以为医护人员及患者提供可靠的安全防护,同时也可用于生物制药、实验检测、食品加工、美容美发、家居清洁、轻工业及电子等行业。



# 产品性能及优点:

- 现有规格: S, M, L;
- 不消毒,一次性使用,左右手均可穿戴;
- 手套呈乳白或淡黄色,双手通用,灵活佩戴;
- 使用进口天然胶乳,柔软舒适、弹性好、不易变形;
- 全麻表面防滑处理,干湿环境均可呈现优越的抓握力;
- 采用真正无粉工艺,使用前可省略掸粉环节,穿戴更方便、更爽滑、更舒适;
- 使用低蛋白天然胶乳,过敏原含量更低,可降低因使用本产品造成的过敏风险。

**包装规格:** ● 100只/盒,盒子尺寸: 24.5x12.5x6.5cm ● 2000只/箱,外箱尺寸: 51x35x27cm \*我们可以按照客户提出的包装要求做成消毒包装。

# 物理性能和尺寸规格参数表 (参照GB10213最新标准)

尺寸代码	标称尺寸	最小长度	宽度	最小厚度	老化前		老化后	
アンコロコ		单位: mm	单位: mm	单位: mm	扯断力 N	伸长率%	扯断力 N	伸长率%
6和6以下	特小号(XS)	220	≤80					
6.5	小号(S)	220	80±5					
7	中号 ( M )	230	85±5	₩ <b>∓</b> 0 00				
7.5	中号 ( M )	230	95±5	光面 0.08 麻面 0.11	≥7.0	≥650	≥6.0	≥500
8	大号 (L)	230	100±5	MACHIOLIT				
8.5	大号 ( L )	230	110±5					
9	特大号(XL)	230	≥110					

销售热线: 010-85307665 - 27 -

# 产品符合欧盟EN455、ISO11193、ASTM D3578、GB10213最新版标准, 并获得欧盟CE证书和美国FDA 510(K)号。

### 第一类医疗器械生产备案凭证

各案编号:	京通食药监械生产各201400044	

企业名称	音樂欄句: 京通資約並做生产各20140004至 名称 北京聯京乳放制品有限公司							
住所	北京市週州区台湖镇北神树村东光机电一体化产业基地兴光五街6号							
生产地址	北京市通州区台湖镇北神树村东光机电一体化产业基地兴光五衡6号							
法定代表人	育文操	企业负责人	削文禄					
生产范围	I类,I-6864医用卫生材料及数料***							
	产品名称	产品各案号	登载日期	备注				
	检查手套	京通城各20140013 号	2014-12-19					
生产产品列表	-							
	-	-						
	-		-					
			-					



CERTIFICAT

TOWSUB TOWSOD TOWSOD TOWSOD TOWSOD T

CERTIFICATE

ZERTIFIKAT







### **EC Certificate**

Production Quality Assurance System
Directive 93/42/EEC on Medical Devices (MDD), Annex V
(Devices in class I in sterile conditions, sterilised systems or procedure packs) No. G2S 044992 0016 Rev. 01

Manufacturer

Beijing Reagent Latex Products Co., Ltd.
Ciqu Industrial Zone, Tongzhou District 101111 Beijing PEOPLE'S REPUBLIC OF CHINA

Facility(ies):

Beijing Reagent Latex Products Co., Ltd. Ciqu Industrial Zone, Tongzhou District, 101111 Beijing, PEOPLE'S REPUBLIC OF CHINA

Product

Sterile Disposable Examination Gloves.

Category(ies):

Valid from: Valid until:

Date, 2019-07-03

1. Pumil

Page 1 of 1 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TUV®

# 第一类医疗器械生产备案证书



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

MAY 2 5 2001

Beijing Reagent Latex Products C/O Ms. Christina Smith Smith Associates P.O. Box Crofton, Maryland 21114

Re: K010947
Trade/Device Name: Snow Lotus Powder Free Latex
Examination Gloves with a Protein Content Labeling
Claim (50 Micrograms or Less)
Regulation Number: 880.6250
Regulatory Class: 1
Product Code: 91
Dated: March 29

Dear Ms. Smith:

Dear Ms. Smith:

We have reviewed your Section 510(k) notification of intent to market the device referenced above and we have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the maximum of the feel of the feel

against misbranding and adulteration.

If your device is classified (see above) into either class II
(Special Controls) or class III (Premarket Approval), it may be
(Special Controls) or class III (Premarket Approval), it may be
control or classified (see above) into a service and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or classified (see a service) and it is a service or complaint or comply with the GMP regulation may result in regulatory action. In addition, FDA may publish further announcements concerning your device in the Federal Register.

## 欧盟CE证书

Page 2 - Ms. Smith

Please note: this response to your premarket notification submission does not affect any obligation you might have under sections 531 through 542 of the Act for devices under the Electronic Product Radiation Control provisions, or other Pederal laws or regulations.

This letter will allow you to begin marketing your device as described in your 510(K) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation [2] CFR part 201 and additionally 80 10 for involved diagnostic devices) please contact the office of Compliance at (301) 594-4522. Additionally, for questions on the promotion and advertising of your device, please contact the Office of Compliance at (301) 594-4639. Also, please note the regulation compliance at (301) 594-4639. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR 807.97). Other general information on your responsibilities under the Act may be obtained from the Division of Small Manufacturera Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its internet address "http://www.fda.gov/cdrh/dsma/dsmamain.html".

Sincerely yours,

Susan Runner

finothy A. Ulatowski
Director
Division of Dental, Infection Control
and General Hospital Devices
Office of Device Svaluation
Center for Devices and
Radiological Health

Enclosure