Note: This report is issued subject to the Testing and Certification Regulations of the TÜV SÜD Group and the General Terms and Conditions of Business of TÜV SÜD PSB Pte Ltd. In addition, this report is governed by the terms set out within this report.

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SUBJECT

Testing of Gloves

CLIENT

Shijiazhuang Hongray Group Co.,Ltd. South Tongda Rd., East Dist., Jinzhou City, Hebei, 052260, China

SAMPLE SUBMISSION DATE/ TEST DATE

29 Mar 2021/01 Apr 2021 to 06 May 2021

DESCRIPTION OF SAMPLES

S/N	Product Description	Brand/ Model	Colour	Lot No.	Size	Sample Received (pieces)	Manufacturer
1	1 Disposable Nitrile Gloves	Hongray Blue		- AP 70.	S	400	
2			- (\$00	М	400	Shijiazhuang	
3			Diue	(See Remark 1)	₋ L	400	Hongray Group Co.,Ltd.
4			-0.000		XL	400	00.,=.a.

METHOD OF TEST:

- 1. EN 455-1:2020 Medical gloves for single use Part 1: Requirements and testing for freedom from holes
- 2. EN 455-2:2015 Medical gloves for single use Part 2: Requirements and testing for physical properties
- 3. EN 455-3:2015 Medical glove for single use Part 3: Requirements and testing for biological evaluation



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RESULTS:

Sample: Disposable Nitrile Gloves, Hongray, Blue

Table 1: Results for EN 455-1:2020

Clause	Tests	Size	Requirements	No. of non- compliers allowed (pieces)	Number tested (pieces)	Actual no. of non- compliers found (pieces)	Inferred results
4 5	Freedom from holes	М	Shall not leak	10	315	2	Passed

Table 2: Results for EN 455-2:2015 Clauses 4-5

Clause	Tests	Size	Requirements (Median)	Number tested (pieces)	Results (Median)	Inferred results
	Dimensions a) Length (mm)	S	≥ 240	13	250	Passed
		М		13	249	Passed
		L		13	246	Passed
4		XL		13	248	Passed
4	b) Width (mm)	S	80 ± 10	13	84	Passed
		М	95 ± 10	13	96	Passed
		L	110 ± 10	13	115	Passed
		XL	≥ 110	13	114	Passed
	Strength a) Force at break (N)	S	For nitrile examination gloves: ≥ 6.0	13	7.5	Passed
		М		13	8.1	Passed
		L		13	8.3	Passed
5		XL		13	7.5	Passed
	b) Force at break after challenge testing (N) 7 days at	S	For nitrile	13	7.6	Passed
		М	M examination gloves:	13	8.1	Passed
		L		13	8.0	Passed
	(70±2)°C	XL		13	7.7	Passed

Table 3: Results for EN 455-2:2015 Clause 7

Clause	Tests	Requirements	Results	Inferred results
7	Labelling	Manufacturers shall label the glove and/or the packaging with the date of manufacture in accordance with EN ISO 15223-1:2012 and EN 1041:2008+A1:2013. Date of manufacture is defined as the packaging date.	Observed	Passed



RESULTS (cont'd):

Sample: Disposable Nitrile Gloves, Hongray, Blue

Table 4: Results for EN 455-3:2015 Clauses 4.2-4.5

Clause	Tests	Requirements	Results / Remarks	Inferred results
4.2	Chemicals	Gloves shall not be dressed with talcum powder (magnesium silicate).	Glove is not dressed with talcum powder, based on client's declaration letter	Passed
		Other chemicals	Manufacturer shall disclose upon request a list of chemical ingredients	NA
4.3 5.1	Endotoxins	< 20 EU/pair for gloves labelled with 'low endotoxin content'.	Not labelled with 'low endotoxin content'	NA
4.4 5.2	Powder- free gloves	For powder-free gloves: The total quantity of powder residues shall not exceed 2 mg per glove.	S 1.24 mg per glove M 0.91 mg per glove L 0.83 mg per glove XL 0.27 mg per glove	Passed Passed Passed Passed
4.5 5.3	Proteins, leachable	The manufacturer shall strive to minimize the leachable protein level for gloves containing natural rubber latex.	Not natural rubber latex glove	NA

Table 5: Results for EN 455-3:2015 Clause 4.6

Clause	Tests	Requirements	Results
	Labelling	In addition to the labelling specified in EN 1041:2008+A1:2013 and the relevant symbols given in EN ISO 15223-1:2012, the following requirements apply:	
		 a) medical gloves containing natural rubber latex shall be labelled on the packaging of at least the smallest packaging unit with the EN ISO 15223-1:2012 symbol for latex; 	NA
		The labelling shall include the following or equivalent warning statement together with the symbol: '(Product) contains natural rubber latex which may cause allergic reactions, including anaphylactic responses';	NA
4.6		 b) the labelling shall include a prominent indication of whether the glove is powdered or powder-free; 	Comply
		 c) sterile powdered gloves shall be labelled with the following or equivalent: 'CAUTION: Surface powder shall be removed aseptically prior to undertaking operative procedures in order to minimize the risk of adverse tissue reactions'; 	NA
		 d) for any medical glove containing natural rubber latex the product labelling shall not include: - any term suggesting relative safety, such as low allergenicity, hypoallergenicity or low protein; - any unjustified indication of the presence of allergens; 	NA
		e) if the manufacturer labels the gloves with the protein content, the process limit, measured as specified in 5.3 shall be given.	NA
		Inferred results	Passed



REMARKS:

- 1. Lot No. was not provided by client.
- 2. Labelling requirements were assessed based on submitted packaging artwork by client.
- 3. NA: Not applicable for the submitted sample.
- 4. The colour as "Blue" was declared by client.

Yeo Poh Kwang Associate Engineer Wong Bee Hui Product Manager Medical Health Services (NAM)

APPENDIX:



Photo 1: Disposable Nitrile Gloves, Hongray, Blue



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Effective 26 January 2021