

LATEX

Comfort-fit

Powder-free Examination Gloves



Recommended
for medical use



- Ideal glove thickness for vigorous procedures
- Cost saving alternative for supporting medical team
- Exceptional elongation & dexterity
- Snug fit ability
- Strong grip
- Renewable resource - natural rubber latex material
- Chemical resistant
- Recyclable packaging

PS54

Reference code: IMPS54Y

PRODUCT SPECIFICATION (ASTM D6319 & EN 455)

Colour: Natural colour

Dimension (mm)	Spec	Typical	
		mm	mil
Cuff	Min. 0.04	0.05	2.0
Palm	Min. 0.05	0.06	2.4
Finger	Min. 0.06	0.08	3.1
Length	Min. 240	240 - 245	

Physical Properties

		Unaged
Tensile strength (MPa)	Min. 18	28 - 32
Elongation (%)	Min. 500	500 - 540
Force at break (N)	Min. 6.0	6.0 - 6.3

Aged

Tensile strength (MPa)	Min. 14	29 - 33
Elongation (%)	Min. 400	460 - 500
Force at break (N)	Min. 6.0	6.0 - 6.3

PACKAGING INFORMATION

Case dimension (mm)	365 x 250 x 250
Box dimension (mm)	240 x 120 x 70
Packing mode	100 pcs / box, 10 boxes / carton

Maximum loading	20'GP	40'GP	40'HQ
	1215 cartons	2600 cartons	3000 cartons

PRODUCT ATTRIBUTES

Medical • Polymer coated • Smooth surface • Standard cuff 9.5" • AQL 1.5



iNtouch™

NITRILE

Shield-aid

Powder-free Examination Gloves



Recommended
for medical use



- Cost saving alternative for supporting medical team
- Excellent film strength
- Excellent puncture resistance
- Customised fitting property, prevent hand fatigue
- Clear indication of tears and breaks
- Prevent type I hypersensitivity
- Chemical resistant
- Recyclable packaging

CS30

Reference code: IMCS30

PRODUCT SPECIFICATION (ASTM D6319 & EN 455)

Colour: Natural colour, blue

Dimension (mm)	Spec	Typical	
		mm	mil
Cuff	Min. 0.04	0.05	2.0
Palm	Min. 0.05	0.06	2.4
Finger	Min. 0.06	0.08	3.1
Length	Min. 240	240 - 245	

Physical Properties

Unaged

Tensile strength (MPa)	Min. 18	28 - 32
Elongation (%)	Min. 500	500 - 540
Force at break (N)	Min. 6.0	6.0 - 6.3

Aged

Tensile strength (MPa)	Min. 14	29 - 33
Elongation (%)	Min. 400	460 - 500
Force at break (N)	Min. 6.0	6.0 - 6.3

PACKAGING INFORMATION

Case dimension (mm)	250 x 240 x 245
Box dimension (mm)	240 x 120 x 45
Packing mode	100 pcs / box, 10 boxes / carton

Maximum loading	20'GP	40'GP	40'HQ
	1860 cartons	3960 cartons	4410 cartons

PRODUCT ATTRIBUTES

Medical • Chlorinated • Fingertips textured • Standard cuff 9.5" • AQL 1.5



iNtouch™

 KOSSAN <small>STRETCHING LIMITS • SINCE 1979</small>	<h1>PRODUCT SPECIFICATION</h1>	Doc No	
		PS-0003	
		Rev	Page
		2	2 of 2

Product : Non-Sterile, Ambidextrous, Finger-Textured Surface, Powder Free Nitrile Examination Gloves

Color : White, Blue, Black

Product Code : OCPFNF-BF-3.0 (EB53001JA-BF), OCPFNF-WC-3.0 (EB53002JA-WC), OCPFNF-BT-3.0 (EB53002JA-BT), OCPFNF-KB-3.0 (EB53001JA-KB)

Reference Code : CS30-BF, CS30-WC, CS30-BT, CS30-KB

Grade : AQL 1.5

Inspection Method : ASTM D6319 and EN 455-1/2/3 (Current Version)

Quality Assurance : Manufactured under ISO 13485, CAN/CSA ISO 13485, ISO 9001 and US FDA QSR Quality Management System

No	Test	Particular	Specification	In-house Inspection Level / AQL	
1	Dimension (mm)	Width	Extra Small	75 ± 5	S2/AQL 4.0
			Small	85 ± 5	
			Medium	95 ± 5	
			Large	106 ± 5	
			Extra Large	116 ± 5	
			Extra Extra Large	125 ± 5	
		Length	Min. 240		
		Thickness (Single Wall)	Cuff (25 ± 5 from bead)	Min. 0.04	
	Palm (center of palm)	Min. 0.05			
	Finger (13 ± 3 from tip)	Min. 0.06			
2	Physical Properties	Unaged (ASTM)	Tensile Strength (MPa)	Min. 18	S2/AQL 4.0
			Elongation at Break (%)	Min. 500	
		Unaged (EN)	Force at Break (N)	Min. 6.0	n = 13 (Median Value)
		Aged (ASTM)	Tensile Strength (MPa)	Min. 14	S2/AQL 4.0
			Elongation at Break (%)	Min. 400	
Aged (EN)	Force at Break (N)	Min. 6.0	n = 13 (Median Value)		
3	Freedom from Holes			GI/AQL 1.5	
4	Visual Defects (Major)	Crack Line, Deformed Gloves, Dirt/Stain ≥1mm ² , Discoloration, Flow Mark, Incomplete Beading, Lump ≥2mm ² , Mix Size, Mix Type, Powder Mark, Polymer Mark, Sticky Gloves/Sticky Pleat ≥5mm, Thin Spot/Fish Eye		GI/AQL 2.5	
5	Visual Defects (Minor)	Blister, Dirt/Stain <1mm ² , Lump <2mm ² , Poor Beading, Sticky Pleat <5mm		GI/AQL 4.0	
6	Powder Residue	Max 1.5 mg/glove		n = 5	

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	<h1>PRODUCT SPECIFICATION</h1>	Doc No	
		PS-1025	
		Rev	Page
		1	2 of 2

Product : Non-Sterile, Ambidextrous, Smooth Surface, Powder Free Polymer Coated, Natural Rubber Latex Examination Gloves

Color : Natural

Product Code : OLPYS-NA-5.4 (ER35401AA-NA)

Reference Code : PS54Y

Grade : AQL 1.5

Inspection Method : ASTM D3578 and EN 455-1/2/3 (Current Version)

Quality Assurance : Manufactured under ISO 13485, CAN/CSA ISO 13485, ISO 9001 and US FDA QSR Quality Management System

No	Test	Particular	Specification	In-house Inspection Level / AQL	
1	Dimension (mm)	Width	Extra Small	75 ± 5	S2/AQL 4.0
			Small	85 ± 5	
			Medium	95 ± 5	
			Large	106 ± 5	
			Extra Large	116 ± 5	
			Extra Extra Large	125 ± 5	
		Length	Min. 240		
		Thickness (Single Wall)	Cuff (25 ± 5 from bead)	Min. 0.06	
Palm (center of palm)	Min. 0.08				
Finger (13 ± 3 from tip)	Min. 0.10				
2	Physical Properties	Unaged (ASTM)	Tensile Strength (MPa)	Min. 18	S2/AQL 4.0
			Elongation at Break (%)	Min. 650	
		Unaged (EN)	Force at Break (N)	Min. 6.0	n = 13 (Median Value)
			Aged (ASTM)	Tensile Strength (MPa)	Min. 14
		Elongation at Break (%)		Min. 500	
Aged (EN)	Force at Break (N)	Min. 6.0	n = 13 (Median Value)		
3	Freedom from Holes			GI/AQL 1.5	
4	Visual Defects (Major)	Crack Line, Deformed Gloves, Dirt/Stain ≥1mm ² , Discoloration, Flow Mark, Incomplete Beading, Lump ≥2mm ² , Mix Size, Mix Type, Powder Mark, Polymer Mark, Sticky Gloves/Sticky Pleat ≥5mm, Thin Spot/Fish Eye		GI/AQL 2.5	
5	Visual Defects (Minor)	Blister, Dirt/Stain <1mm ² , Lump <2mm ² , Poor Beading, Sticky Pleat <5mm		GI/AQL 4.0	
6	Powder Residue	Max 1.5 mg/glove		n = 5	
7	Protein	Max 50 µg/g (for EN Testing Method)		n = 8	
		Max 50 µg/dm ² (for ASTM Testing Method)		n = 3	

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Digitally signed by Dicusar Vladimir
Date: 2022.09.21 17:19:41 EEST
Reason: MoldSign Signature
Location: Moldova



• Powder-free Examination Gloves
• Chemical permeation tested • Viral penetration tested • Fentanyl tested • Chemotherapy drugs tested

NITRILE
Shield-aid

XS
5 - 6
100 gloves
by weight

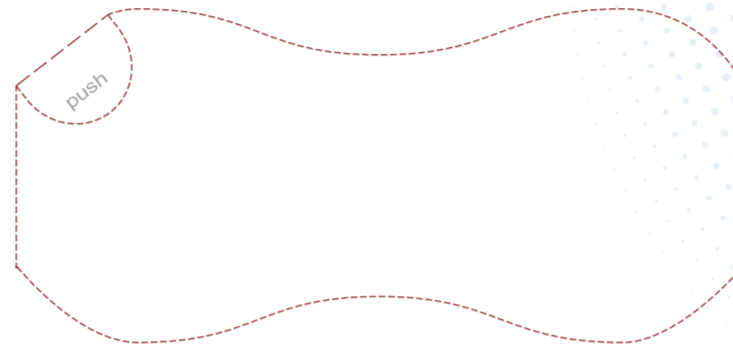
NITRILE LD
Shield-aid
Powder-free Examination Gloves

NITRILE

Shield-aid

CS30

Powder-free Examination Gloves



BLUE
XS
5 - 6
100 gloves
by weight



Recommended for medical use

BLUE
XS
5 - 6
100 gloves
by weight

intouch
TM

BLUE
XS
5 - 6
100 gloves
by weight

NITRILE LD
Shield-aid
Powder-free Examination Gloves

XS

NITRILE

Shield-aid

Powder-free Examination Gloves

- Gants d'examen nitrile sans poudre
- Puderfrei untersuchun gshandschuhe aus nitril
- Mănuși nitril nepudrate pentru examinare

XS
5 - 6
100 gloves
by weight

XS

NITRILE

Shield-aid

Powder-free Examination Gloves

In compliance with Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III). Notified Body responsible for PPE EU Type Examination (Module B) and Module C2 On-going Conformity Assessment is SATRA Technology Europe Limited (2777) Bracetown Business park, Clonee, Dublin 15, D15 YN2P, Ireland. EU Declaration of Conformity is accessible at www.intouchcares.com. Product reference: PFN

In compliance with Medical Devices Regulations 2002 (SI 2002 No. 618, as amended) (UK MDR 2002) and PPE Regulation (EU) 2016/425 as retained in UK Law and amended. UK Approved Body responsible for PPE UKCA Type Examination (Module B) and Module C2 On-going Conformity Assessment is SATRA Technology Center Limited (AB0521), Wyndham Way, Kettering NN16 8SD, United Kingdom. In compliance with PPE Regulation (EU) 2016/425 as retained in UK Law and amended. UK Declaration of Conformity is accessible at www.intouchcares.com.

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User information sheet is enclosed in this package.

KOSSAN INTERNATIONAL SDN. BHD. (2731784M) Tel +603 3392 3013
Wisma KOSSAN, Lot 782, Jalan Sungai Putus, Off Batu 3rd, Jalan Kapar, 42100 Klang, Selangor, Malaysia.

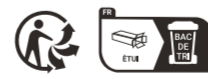
ADVENA LIMITED Email info@advena.com
Tower Business Centre, 2nd Floor, Tower Street, Swatara, BKR 4013, Malta.

UKRP ADVENA LIMITED UK Email info@advenamedical.com
Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

Made in Malaysia



REF
IMCS30BF-XS



Tested for use with chemotherapy drugs using ASTM D6978.

Chemotherapy drugs & chemicals	Breakthrough detection time
Cisplatin 1 mg/ml, Cyclophosphamide (Cytoxan) 20 mg/ml, Cytarabine 100 mg/ml, Docetaxel (Taxot) 10 mg/ml, Doxorubicin hydrochloride 2 mg/ml, Etoposide 20 mg/ml, Fluorouracil 50 mg/ml, Ifosfamide 50 mg/ml, Methotrexate 25 mg/ml, Mitomycin C 0.5 mg/ml, Mitoxantrone 2 mg/ml, Paclitaxel (Taxol) 6 mg/ml, Vincristine sulfate 1 mg/ml	Minimum 240 min
Thiotepa 10 mg/ml	30.2 min
Carmustine (BCNU) 3.3 mg/ml	10.1 min
Fentanyl citrate injection (100 µg/2ml)	Minimum 240 min

Warning: Not recommended for use with Carmustine and Thiotepa.

Chemicals	Level	Mean degradation	Chemicals	Level	Mean degradation
*4% Chlorhexidine Digluconate	6	19.0%	10% Sodium Percarbonate	6	15.4%
40% Sodium Hydroxide (K)	6	42.9%	10% Acetic Acid	4	66.7%
10-13% Sodium Hypochlorite	6	14.7%	37% Formaldehyde (T)	3	5.0%
50% Sulphuric Acid	6	20.5%	30% Hydrogen Peroxide (P)	2	22.8%
5% Ethidium Bromide	6	3.4%	25% Ammonium Hydroxide (O)	0	-52.0%
50% Glutaraldehyde	6	27.4%	65% Nitric Acid (M)	0	97.6%
0.1% Phenol	6	33.8%	70% Isopropanol	0	62.2%
1.5% Methanol in Water	6	21.9%	35% Ethanol	0	38.8%
3% Povidone-iodine	6	33.7%	99% Acetic Acid (N)	0	93.9%

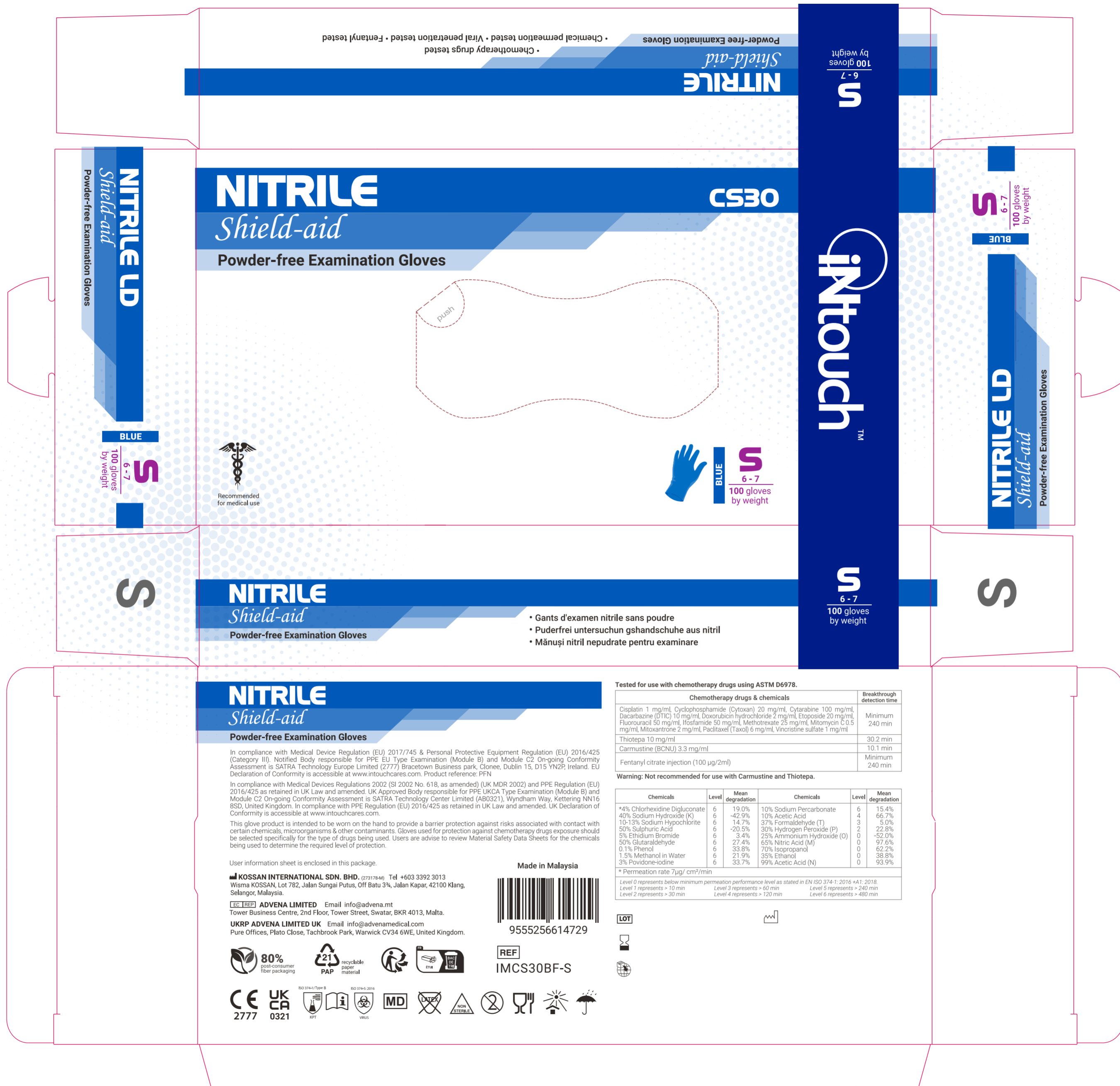
* Permeation rate 7 µg/cm²/min

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018.
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min

LOT



235 x 120 x 45



• Chemotherapy drugs tested
• Viral penetration tested
• Fentanyl tested

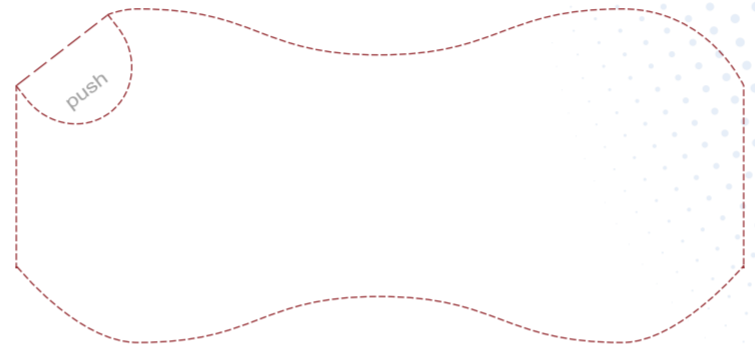
NITRILE
Shield-aid

S
6-7
100 gloves
by weight

NITRILE
Shield-aid

CS30

Powder-free Examination Gloves



NITRILE LD
Shield-aid
Powder-free Examination Gloves

BLUE
100 gloves
by weight
S
6-7



BLUE
100 gloves
by weight
S
6-7

S
6-7
100 gloves
by weight
BLUE

NITRILE LD
Shield-aid
Powder-free Examination Gloves

S

NITRILE
Shield-aid
Powder-free Examination Gloves

- Gants d'examen nitrile sans poudre
- Puderfrei untersuchungshandschuhe aus nitril
- Mănuși nitril nepudrate pentru examinare

S
6-7
100 gloves
by weight

S

NITRILE
Shield-aid
Powder-free Examination Gloves

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UKRP ADVENA LIMITED UK Email info@advenamedical.com
Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

Made in Malaysia



REF
IMCS30BF-S



Tested for use with chemotherapy drugs using ASTM D6978.

Chemotherapy drugs & chemicals	Breakthrough detection time
Cisplatin 1 mg/ml, Cyclophosphamide (Cytoxan) 20 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10 mg/ml, Doxorubicin hydrochloride 2 mg/ml, Etoposide 20 mg/ml, Fluorouracil 50 mg/ml, Ifosfamide 50 mg/ml, Methotrexate 25 mg/ml, Mitomycin C 0.5 mg/ml, Mitoxantrone 2 mg/ml, Paclitaxel (Taxol) 6 mg/ml, Vincristine sulfate 1 mg/ml	Minimum 240 min
Thiotepa 10 mg/ml	30.2 min
Carmustine (BCNU) 3.3 mg/ml	10.1 min
Fentanyl citrate injection (100 µg/2ml)	Minimum 240 min

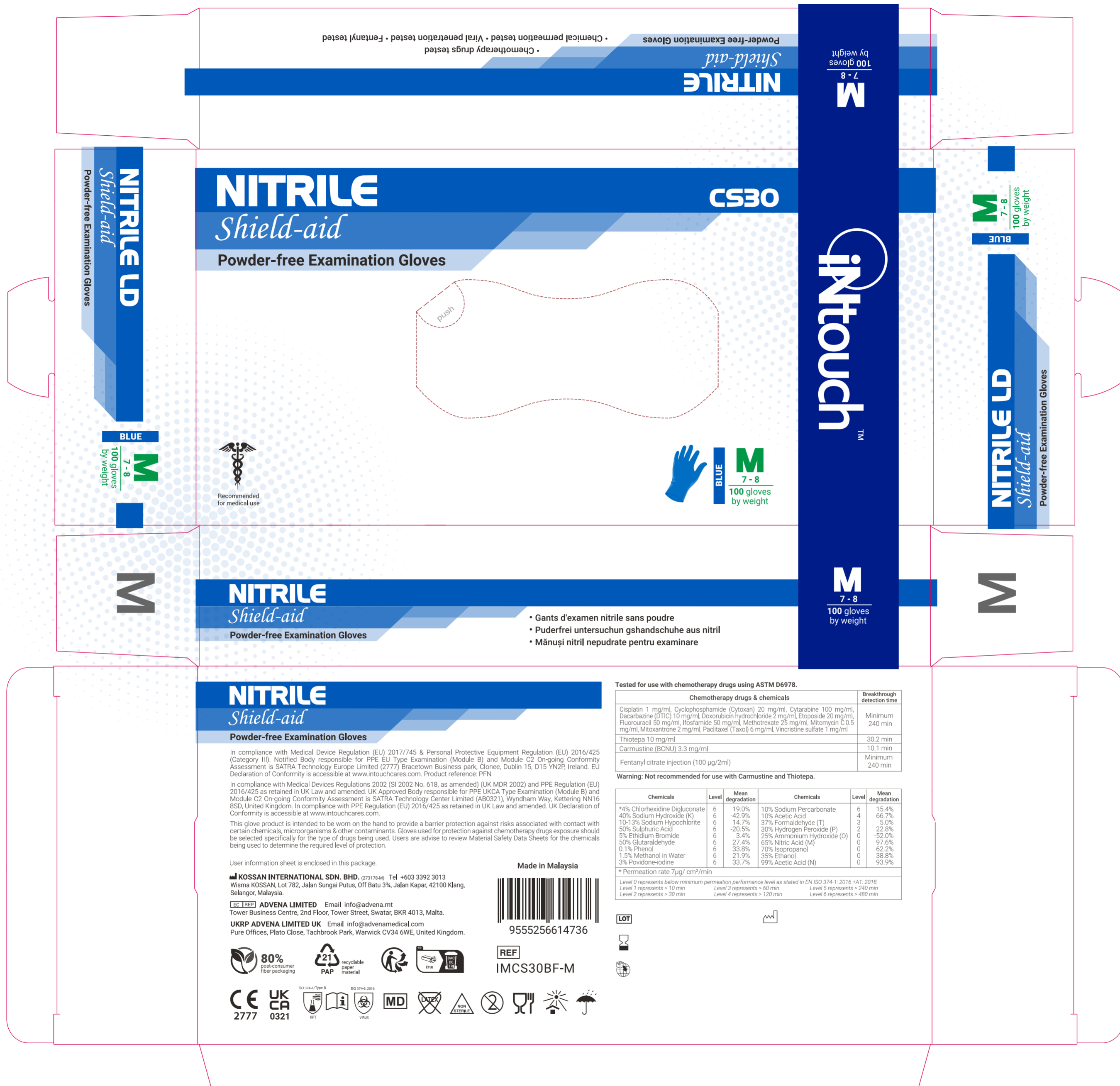
Warning: Not recommended for use with Carmustine and Thiotepa.

Chemicals	Level	Mean degradation	Chemicals	Level	Mean degradation
*4% Chlorhexidine Digluconate	6	19.0%	10% Sodium Percarbonate	6	15.4%
40% Sodium Hydroxide (K)	6	-42.9%	10% Acetic Acid	4	66.7%
10-13% Sodium Hypochlorite	6	14.7%	37% Formaldehyde (T)	3	5.0%
50% Sulphuric Acid	6	-20.5%	30% Hydrogen Peroxide (P)	2	22.8%
5% Ethidium Bromide	6	3.4%	25% Ammonium Hydroxide (O)	0	-52.0%
50% Glutaraldehyde	6	27.4%	65% Nitric Acid (M)	0	97.6%
0.1% Phenol	6	33.8%	70% Isopropanol	0	62.2%
1.5% Methanol in Water	6	21.9%	35% Ethanol	0	38.8%
3% Povidone-iodine	6	33.7%	99% Acetic Acid (N)	0	93.9%

* Permeation rate 7 µg/cm²/min
Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018.
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min



235 x 120 x 45



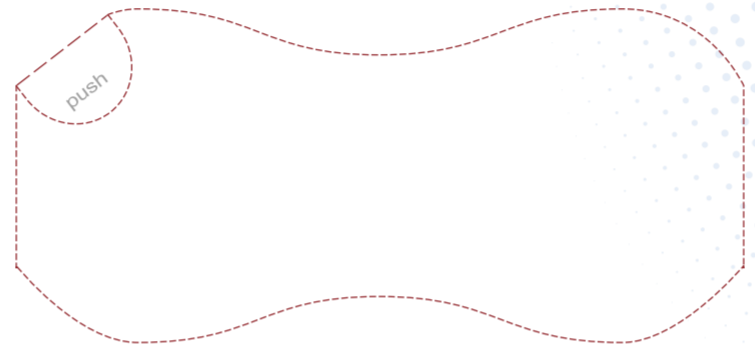
• Powder-free Examination Gloves
 • Chemotherapy drugs tested
 • Viral penetration tested
 • Fentanyl tested

NITRILE
Shield-aid

M
 7-8
 100 gloves
 by weight

NITRILE
Shield-aid
CS30

Powder-free Examination Gloves



NITRILE LD
Shield-aid
 Powder-free Examination Gloves

BLUE
M
 7-8
 100 gloves
 by weight



BLUE
M
 7-8
 100 gloves
 by weight

BLUE
M
 7-8
 100 gloves
 by weight

NITRILE LD
Shield-aid
 Powder-free Examination Gloves

intouchTM

M

NITRILE
Shield-aid
Powder-free Examination Gloves

- Gants d'examen nitrile sans poudre
- Puderfrei untersuchungshandschuhe aus nitril
- Mănuși nitril nepudrate pentru examinare

M
 7-8
 100 gloves
 by weight

M

NITRILE
Shield-aid
Powder-free Examination Gloves

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 Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

Made in Malaysia



REF
IMCS30BF-M



Tested for use with chemotherapy drugs using ASTM D6978.

Chemotherapy drugs & chemicals	Breakthrough detection time
Cisplatin 1 mg/ml, Cyclophosphamide (Cytoxan) 20 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10 mg/ml, Doxorubicin hydrochloride 2 mg/ml, Etoposide 20 mg/ml, Fluorouracil 50 mg/ml, Ifosfamide 50 mg/ml, Methotrexate 25 mg/ml, Mitomycin C 0.5 mg/ml, Mitoxantrone 2 mg/ml, Paclitaxel (Taxol) 6 mg/ml, Vincristine sulfate 1 mg/ml	Minimum 240 min
Thiotepa 10 mg/ml	30.2 min
Carmustine (BCNU) 3.3 mg/ml	10.1 min
Fentanyl citrate injection (100 µg/2ml)	Minimum 240 min

Warning: Not recommended for use with Carmustine and Thiotepa.

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40% Sodium Hydroxide (K)	6	-42.9%	10% Acetic Acid	4	66.7%
10-13% Sodium Hypochlorite	6	14.7%	37% Formaldehyde (T)	3	5.0%
50% Sulphuric Acid	6	-20.5%	30% Hydrogen Peroxide (P)	2	22.8%
5% Ethidium Bromide	6	3.4%	25% Ammonium Hydroxide (O)	0	-52.0%
50% Glutaraldehyde	6	27.4%	65% Nitric Acid (M)	0	97.6%
0.1% Phenol	6	33.8%	70% Isopropanol	0	62.2%
1.5% Methanol in Water	6	21.9%	35% Ethanol	0	38.8%
3% Povidone-iodine	6	33.7%	99% Acetic Acid (N)	0	93.9%

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235 x 120 x 45

• Chemotherapy drugs tested • Viral penetration tested • Fentanyl tested

NITRILE

Shield-aid

100 gloves
by weight
6 - 8

NITRILE
Shield-aid

CS30

Powder-free Examination Gloves

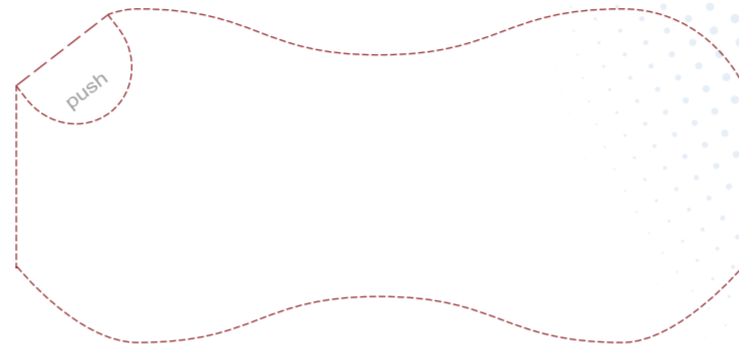
100 gloves
by weight
8 - 9
BLUE

NITRILE LD
Shield-aid
Powder-free Examination Gloves

BLUE
100 gloves
by weight
8 - 9



Recommended for medical use



BLUE
100 gloves
by weight
8 - 9

NITRILE LD
Shield-aid
Powder-free Examination Gloves

intouchTM

8 - 9
100 gloves
by weight

NITRILE
Shield-aid

Powder-free Examination Gloves

- Gants d'examen nitrile sans poudre
- Puderfrei untersuchun gshandschuhe aus nitril
- Mănuși nitril nepudrate pentru examinare

NITRILE
Shield-aid

Powder-free Examination Gloves

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ADVENA LIMITED Email info@advena.com
Tower Business Centre, 2nd Floor, Tower Street, Swatar, BKR 4013, Malta.

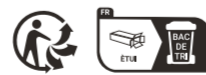
UKRP ADVENA LIMITED UK Email info@advenamedical.com
Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

Made in Malaysia



9555256614743

REF
IMCS30BF-L



Tested for use with chemotherapy drugs using ASTM D6978.

Chemotherapy drugs & chemicals	Breakthrough detection time
Cisplatin 1 mg/ml, Cyclophosphamide (Cytoxan) 20 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10 mg/ml, Doxorubicin hydrochloride 2 mg/ml, Etoposide 20 mg/ml, Fluorouracil 50 mg/ml, Ifosfamide 50 mg/ml, Methotrexate 25 mg/ml, Mitomycin C 0.5 mg/ml, Mitoxantrone 2 mg/ml, Paclitaxel (Taxol) 6 mg/ml, Vincristine sulfate 1 mg/ml	Minimum 240 min
Thiotepa 10 mg/ml	30.2 min
Carmustine (BCNU) 3.3 mg/ml	10.1 min
Fentanyl citrate injection (100 µg/2ml)	Minimum 240 min

Warning: Not recommended for use with Carmustine and Thiotepa.

Chemicals	Level	Mean degradation	Chemicals	Level	Mean degradation
*4% Chlorhexidine Digluconate	6	19.0%	10% Sodium Percarbonate	6	15.4%
40% Sodium Hydroxide (K)	6	-42.9%	10% Acetic Acid	4	66.7%
10-13% Sodium Hypochlorite	6	14.7%	37% Formaldehyde (T)	3	5.0%
50% Sulphuric Acid	6	-20.5%	30% Hydrogen Peroxide (P)	2	22.8%
5% Ethidium Bromide	6	3.4%	25% Ammonium Hydroxide (O)	0	-52.0%
50% Glutaraldehyde	6	27.4%	65% Nitric Acid (M)	0	97.6%
0.1% Phenol	6	33.8%	70% Isopropanol	0	62.2%
1.5% Methanol in Water	6	21.9%	35% Ethanol	0	38.8%
3% Povidone-iodine	6	33.7%	99% Acetic Acid (N)	0	93.9%

* Permeation rate 7µg/cm²/min

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1:2016 +A1:2018
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min

LOT



235 x 120 x 45

• Chemotherapy drugs tested
• Viral penetration tested • Fentanyl tested
• Powder-free Examination Gloves

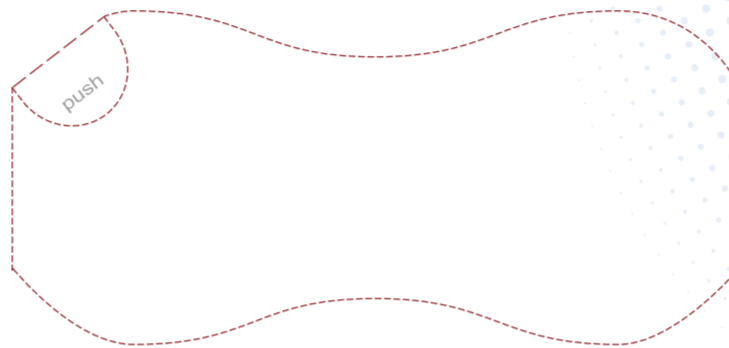
NITRILE
Shield-aid

XL
9 - 10
100 gloves
by weight

NITRILE
Shield-aid

CS30

Powder-free Examination Gloves



NITRILE LD
Shield-aid
Powder-free Examination Gloves

XL
9 - 10
100 gloves
by weight

XL
9 - 10
100 gloves
by weight

NITRILE LD
Shield-aid
Powder-free Examination Gloves

intouchTM

XL
9 - 10
100 gloves
by weight

XL

NITRILE
Shield-aid

Powder-free Examination Gloves

- Gants d'examen nitrile sans poudre
- Puderfrei untersuchungshandschuhe aus nitril
- Mănuși nitril nepudrate pentru examinare

NITRILE
Shield-aid

Powder-free Examination Gloves

In compliance with Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III). Notified Body responsible for PPE EU Type Examination (Module B) and Module C2 On-going Conformity Assessment is SATRA Technology Europe Limited (2777) Bracetown Business park, Clonee, Dublin 15, D15 YN2P, Ireland. EU Declaration of Conformity is accessible at www.intouchcares.com. Product reference: PFN

In compliance with Medical Devices Regulations 2002 (SI 2002 No. 618, as amended) (UK MDR 2002) and PPE Regulation (EU) 2016/425 as retained in UK Law and amended. UK Approved Body responsible for PPE UKCA Type Examination (Module B) and Module C2 On-going Conformity Assessment is SATRA Technology Center Limited (AB0521), Wyndham Way, Kettering NN16 8SD, United Kingdom. In compliance with PPE Regulation (EU) 2016/425 as retained in UK Law and amended. UK Declaration of Conformity is accessible at www.intouchcares.com.

This glove product is intended to be worn on the hand to provide a barrier protection against risks associated with contact with certain chemicals, microorganisms & other contaminants. Gloves used for protection against chemotherapy drugs exposure should be selected specifically for the type of drugs being used. Users are advised to review Material Safety Data Sheets for the chemicals being used to determine the required level of protection.

User information sheet is enclosed in this package.

Made in Malaysia

KOSSAN INTERNATIONAL SDN. BHD. (2731784M) Tel +603 3392 3013
Wisma KOSSAN, Lot 782, Jalan Sungai Putus, Off Batu 3rd, Jalan Kapar, 42100 Klang, Selangor, Malaysia.

ADVENA LIMITED Email info@advena.com
Tower Business Centre, 2nd Floor, Tower Street, Swatar, BKR 4013, Malta.

UKRP ADVENA LIMITED UK Email info@advenamedical.com
Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.



9555256614750

REF
IMCS30BF-XL



Tested for use with chemotherapy drugs using ASTM D6978.

Chemotherapy drugs & chemicals	Breakthrough detection time
Cisplatin 1 mg/ml, Cyclophosphamide (Cytoxan) 20 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10 mg/ml, Doxorubicin hydrochloride 2 mg/ml, Etoposide 20 mg/ml, Fluorouracil 50 mg/ml, Ifosfamide 50 mg/ml, Methotrexate 25 mg/ml, Mitomycin C 0.5 mg/ml, Mitoxantrone 2 mg/ml, Paclitaxel (Taxol) 6 mg/ml, Vincristine sulfate 1 mg/ml	Minimum 240 min
Thiotepa 10 mg/ml	30.2 min
Carmustine (BCNU) 3.3 mg/ml	10.1 min
Fentanyl citrate injection (100 µg/2ml)	Minimum 240 min

Warning: Not recommended for use with Carmustine and Thiotepa.

Chemicals	Level	Mean degradation	Chemicals	Level	Mean degradation
*4% Chlorhexidine Digluconate	6	19.0%	10% Sodium Percarbonate	6	15.4%
40% Sodium Hydroxide (K)	6	-42.9%	10% Acetic Acid	4	66.7%
10-13% Sodium Hypochlorite	6	14.7%	37% Formaldehyde (T)	3	5.0%
50% Sulphuric Acid	6	-20.5%	30% Hydrogen Peroxide (P)	2	22.8%
5% Ethidium Bromide	6	3.4%	25% Ammonium Hydroxide (O)	0	-52.0%
50% Glutaraldehyde	6	27.4%	65% Nitric Acid (M)	0	97.6%
0.1% Phenol	6	33.8%	70% Isopropanol	0	62.2%
1.5% Methanol in Water	6	21.9%	35% Ethanol	0	38.8%
3% Povidone-iodine	6	33.7%	99% Acetic Acid (N)	0	93.9%

* Permeation rate 7 µg/cm²/min

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1:2016 +A1:2018
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min

LOT



235 x 120 x 45

intouch™

POWDER FREE NITRILE EXAMINATION GLOVES

EN : USER INFORMATION
FR : INFORMATIONS DE L'UTILISATEUR
RO : INFORMAȚII UTILIZATOR

Available Size



ISO 374-1/Type B

ISO 374-5:2016



EN POWDER FREE NITRILE EXAMINATION GLOVES

• Non sterile • Single Use
• Recommended for medical use

Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III) • Medical Devices Regulations 2002 (SI 2002 No. 618, as amended) (UK MDR 2002) • EU Notified Body for (Module B) and Module C2: SATRA Technology Europe Limited (2777) • UK Approved Body for (Module B) and Module C2: SATRA Technology Center Limited (A80321) • EU Declaration of Conformity & UK Declaration of Conformity is accessible at www.intouchcares.com. This information does not reflect the actual duration of protection in the workplace and the differentiation between mixtures and pure chemicals. • The chemical resistance has been assessed under laboratory conditions from samples taken from the palm only (except in cases where the glove is equal to or over 400mm – where the cuff is tested also) and relates only to the chemical tested. It can be different if the chemical used in a mixture. • It is recommended to check that the gloves are suitable for the intended use because the conditions at the workplace may differ from the type test depending on temperature, abrasion and degradation. • When used, protective glove may provide less resistance to the dangerous chemical due to changes in physical properties. Movements, snagging, rubbing, degradation caused by chemical contact etc. may reduce the actual use time significantly. For corrosive chemicals, degradation can be the most important factor to consider in selection of chemical resistant gloves. • Before usage, inspect the gloves for any defect or imperfections. • EN ISO 374-4:2019 Degradation levels indicate the change in puncture resistance of the gloves after exposure to the challenge chemical. • The penetration resistance has been assessed under laboratory conditions and relates only to the tested specimens. • Donning – Hold glove by the bead with one hand. Align the glove thumb with your other hand thumb and slide your hand into the glove, one finger into each glove finger. Pull by the glove palm to get a good fit. Don the other glove by the same procedure. • Doffing – Hold glove bead and pull toward the finger until the glove come off. • Where relevant, a list of the substances contained in the glove which are known to cause allergies, per listed in Annex G of EN ISO 21420:2020, shall be supplied on request. • Components used in glove manufacturing may cause allergic reactions in some users. If allergic reactions occur, seek for medical advice immediately. • In storage, avoid excessive heat. Open box should be shielded from exposure to direct sun or fluorescent lighting. Gloves are packed in dispenser which is suitable for transport. Keep the gloves in box when not in use.

Minimum Breakthrough Time (min) >240min	Chemotherapy Drug & Concentration
	Cisplatin 1.0 mg/ml, Cyclophosphamide (Cytoxan) 20.0 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10.0 mg/ml, Doxorubicin Hydrochloride 2.0 mg/ml, Etoposide 20.0 mg/ml, Fluorouracil 50.0 mg/ml, Ifosfamide 50.0 mg/ml, Methotrexate 25.0 mg/ml, Mitomycin 0.5 mg/ml, Mitoxantrone 2.0 mg/ml, Paclitaxel (Taxol) 6.0 mg/ml, Vincristine Sulfate 1.0 mg/ml
	Fentanyl citrate and concentration
	Fentanyl citrate injection (100 mcg/2ml)

30.2min Thiotepa (10.0 mg/ml)
10.1min Carmustine (BCNU) (3.3 mg/ml)

Warning: Not recommended for use with Carmustine and Thiotepa.
Tested for use with Chemotherapy Drugs using ASTM D6978.

Resistance to Permeation by Chemicals	Level	Mean Degradation (%)
*4% Chlorhexidine Digluconate	6	19.0
40% Sodium Hydroxide (K)	6	-42.9
10-13% Sodium Hypochlorite	6	14.7
50% Sulphuric Acid	6	-20.5
5% Ethidium Bromide	6	3.4
50% Glutaraldehyde	6	27.4
0.1% Phenol	6	33.8
1.5% Methanol in Water	6	21.9
3% Povidone-iodine	6	33.7
10% Sodium Percarbonate	6	15.4
10% Acetic Acid	4	66.7
37% Formaldehyde (T)	3	5.0
30% Hydrogen Peroxide (P)	2	22.8
25% Ammonium Hydroxide (O)	0	-52.0
65% Nitric Acid (M)	0	97.6
70% Isopropanol	0	62.2
35% Ethanol	0	93.9
99% Acetic Acid (N)	0	93.9

* Permeation rate 7µg/cm²/min

Permeation Performance Level & Measured Breakthrough Time (minutes)
0 > * 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indicates that the glove falls below the minimum performance level as stated in EN ISO 374-1:2016 +A1:2018 for the given individual hazard.



PAP
RECYCLABLE
PAPER MATERIAL
G-LF-XXXXXX-F-R0
Last update in 2022

FR POWDER FREE NITRILE EXAMINATION GLOVES

• Non sterile • Single Use
• Recommended for medical use

Règlement sur les dispositifs médicaux (UE) 2017/745 et règlement sur les équipements de protection individuelle (UE) 2016/425 (catégorie III). • Règlement sur les dispositifs médicaux de 2002 (SI 2002 No. 618, tel que modifié) (UK MDR 2002) • Organisme notifié de l'UE pour le (module B) et le module C2, est SATRA Technology Europe Limited (2777) • Organisme agréé Royaume-Uni pour le (module B) et le module C2 est SATRA Technology Center Limited (A80321). • La déclaration de conformité de l'UE et la déclaration de conformité du Royaume-Uni sont accessibles à l'adresse www.intouchcares.com. Ces informations ne reflètent pas la durée réelle de la protection sur le lieu de travail et la différenciation entre les mélanges et les produits chimiques purs. • La résistance chimique a été évaluée dans des conditions de laboratoire à partir d'échantillons prélevés sur la paume uniquement (sauf dans les cas où le gant mesure 400 mm ou plus – ou la manchette est également testée) et ne concerne que le produit chimique testé. Elle peut être différente si le produit chimique est utilisé dans un mélange. • Il est recommandé de vérifier que les gants sont adaptés à l'utilisation prévue car les conditions sur le lieu de travail peuvent différer de l'essai de type en fonction de la température, de l'abrasion et de la dégradation. • Lorsque est utilisé le gant de protection peut offrir une moindre résistance au produit chimique dangereux en raison des modifications de ses propriétés physiques. Les mouvements, les accrochages, les frottements, la dégradation causés par le contact chimique, etc. peuvent réduire considérablement la durée d'utilisation réelle. Pour les produits chimiques corrosifs, la dégradation peut être le facteur le plus important à prendre en compte dans le choix de gant de protection aux produits chimiques. • Avant l'utilisation, inspectez les gants pour détecter tout défaut ou imperfection. • EN ISO 374-4:2019 Les niveaux de dégradation indiquent le changement de la résistance à la perforation des gants après exposition au produit chimique de référence. • La résistance à la pénétration a été évaluée dans des conditions de laboratoire et ne concerne que les spécimens testés. • Enfilage - Tenez le gant par le talon d'une main. Alignez le pouce du gant avec le pouce de l'autre main et glissez votre main dans le gant, un doigt dans chaque doigt et tirez sur la paume du gant pour obtenir un bon ajustement. Enfilez l'autre gant en suivant la même procédure. • Enlever le gant - Tenir le talon du gant et tirer vers le doigt jusqu'à ce que le gant se détache. • Le cas échéant, une liste des substances contenues dans le gant et connues pour provoquer des allergies, telles qu'énumérées à l'annexe G de la norme EN ISO 21420:2020, doit être fournie sur demande. • Les composants utilisés dans la fabrication des gants peuvent provoquer des réactions allergiques chez certains utilisateurs. En cas de réaction allergique, consulter immédiatement un médecin. Lors du stockage, éviter toute chaleur excessive. La boîte ouverte doit être protégée de l'exposition directe au soleil ou à un éclairage fluorescent. Les gants sont emballés dans un distributeur qui convient au transport. Conservez les gants dans leur boîte lorsqu'ils ne sont pas utilisés.

Temps de détection minimal de la percée > 240min	Médicament de chimiothérapie et concentration
	Cisplatine 1,0 mg/ml, Cyclophosphamide (Cytoxan) 20,0 mg/ml, Cytarabine 100 mg/ml, Dacarbazine (DTIC) 10,0 mg/ml, Doxorubicin Hydrochloride 2,0 mg/ml, Etoposide 20,0 mg/ml, Fluorouracil 50,0 mg/ml, Ifosfamide 50,0 mg/ml, Methotrexate 25,0 mg/ml, Mitomycine 0,5 mg/ml, Mitoxantrone 2,0 mg/ml, Paclitaxel (Taxol) 6,0 mg/ml, Sulfate de Vincristine 1,0 mg/ml
	Citrate de fentanyl et concentration
	Injection de citrate de fentanyl (100 mcg/2ml)

30.2min Thiotepa (10.0 mg/ml)
10.1min Carmustine (BCNU) (3.3 mg/ml).

avertissement : Non recommandé pour une utilisation avec Carmustine et Thiotepa.
Testé pour une utilisation avec les médicaments de chimiothérapie selon ASTM D6978.

Résistance à la perméation par les produits chimiques	Niveau	Moyenne Dégradation (%)
*4% Digluconate de chlorhexidine	6	19,0
Hydroxyde de sodium à 40% (K)	6	-42,9
10-13% Hypochlorite de sodium	6	14,7
50% Acide sulfurique	6	-20,5
5% Bromure d'éthidium	6	3,4
50% Glutaraldéhyde	6	27,4
0,1% Phénol	6	33,8
1,5% Méthanol dans l'eau	6	21,9
3% Povidone-iodée	6	33,7
10% Carbonate de sodium	6	15,4
10% Acide acétique	4	66,7
37% Formaldéhyde (T)	3	5,0
30% Peroxyde d'hydrogène (P)	2	22,8
25% D'hydroxyde d'ammonium (O)	0	-52,0
65% Acide nitrique (M)	0	97,6
70% Isopropanol	0	62,2
35% Ethanol	0	93,9
99% Acide acétique à 99 % (N)	0	93,9

* Taux de perméation 7µg/cm²/min

Niveau de performance de perméation et temps de percée mesurés (minutes)
0 > * 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indique que le gant est inférieur au niveau de performance minimum tel qu'indiqué dans la norme EN ISO 374-1:2016 +A1:2018 pour le risque individuel donné.

RO POWDER FREE NITRILE EXAMINATION GLOVES

• Non sterile • Single Use
• Recommended for medical use

Regulamentul privind Dispozitivele Medicale (UE) 2017/745 și Regulamentul privind Echipamentul Individual de Protecție (UE) 2016/425 (Categorie III). • Regulamentul privind Dispozitivele Medicale 2002 (SI 2002 Nr. 618, cu modificări) (UK MDR 2002) • Organismul Notificat UE pentru (Modulul B) și Modulul C2: SATRA Technology Europe Limited (2777) • Organismul Notificat MB pentru (Modulul B) și Modulul C2: SATRA Technology Center Limited (A80321) • Declarația de Conformitate UE și Declarația de Conformitate MB este accesibilă la adresa www.intouchcares.com. Informația dată nu se referă la protecția locului de muncă și siguranța la diferite amestecuri din produse chimice. • Rezistența chimică a fost evaluată în condiții de laborator, mostrele folosite au fost selectate doar din palmă, cu excepția cazurilor când mărșă are 400 mm și mai mult – unde și manșeta este testată) și se referă doar la substanța chimică testată. Rezultatele pot fi diferite dacă este folosit un amestec de substanțe chimice. • Se recomandă de a verifica dacă mărșă sunt potrivite pentru a fi folosite în scopul propus, deoarece condițiile locului de muncă pot fi diferite de tipul testărilor efectuate și se pot diferența prin temperatură, abraziune și degradare. • Mărșăle de protecție, în timpul utilizării, pot oferi o rezistență mai mică la substanțele chimice periculoase din cauza modificărilor proprietăților fizice la acțiunea acestor compuși chimici. Mișcările, strângerea, frecarea, degradarea cauzată de interacțiunea chimică etc. poate reduce semnificativ timpul real de utilizare. Pentru produsele chimice corozive, degradarea poate fi cel mai important factor pentru a fi luat în considerare la selectarea mărșăle rezistente la produsele chimice. • Înainte de folosire, verificați mărșăle la lipsa defectelor și imperfecțiunilor. • EN ISO 374-4:2019 Nivelurile de degradare indică modificarea rezistenței mărșăle la perforare după expunerea la substanțele chimice. • Rezistența la penetrare a fost evaluată în baza condițiilor de laborator și se referă doar la specișimenele testate. • Imbrăcarea - Așezați mărșă de margine cu o mână. Introduceți degetul mare al mărșăle pe degetul mare al celeilalte mărșă și trageți mărșă în mărșă, fiecare deget al mărșăle în degetul mărșăle. Trageți astfel mărșăle încât să se potrivească. Imbrăcați celelalte mărșă după același scenariu. • Dezbrăcarea - Țineți marginea mărșăle și trageți de pe degete până mărșăle va iesi. • Unde e cazul, se va oferi la cerere lista componentelor mărșăle care sunt recunoscute ca alergeni, menționate în Anexa G a EN ISO 21420:2020. • Componentele folosite la producerea mărșăle pot cauza reacții alergice la unii utilizatori. Dacă se produce o reacție alergică, adresată-vă urgent medicului. • Mărșăle se vor păstia la loc ferit de căldură în exces. Cătu deschisă se va proteja de expunerea directă la soare sau lumină fluorescentă. • Mărșăle sunt ambalate în cutie care se potrivește pentru transportare. Păstrați mărșăle în cutie atunci când nu sunt folosite.

Temp minim de detectare a perforării > 240min	Medicamente chimioterapice și concentrație
	Cisplatina 1,0 mg/ml, Ciclofosfamidă (Cytoxan) 20,0 mg/ml, Citarabină 100 mg/ml, Dacarbazină (DTIC) 10,0 mg/ml, Doxorubicină clorhidrat 2,0 mg/ml, Etoposid 20,0 mg/ml, Ifos.0 mg/ml, Ios.0 mg/ml, Metotrexat 25,0 mg/ml, Mitomicina 0,5 mg/ml, Mitoxantrona 2,0 mg/ml, Paclitaxel (Taxol) 6,0 mg/ml Sulfat de vincristina 1,0 mg/ml
	Citrat de fentanil și concentrație
	Injecție cu citrat de fentanil (100 mcg/2ml)

30.2min Thiotepa (10.0 mg/ml)
10.1min Carmustină (BCNU) (3.3 mg/ml).

Atenție: Nu se recomandă folosirea cu Carmustină și Thiotepa.
Testate la folosirea cu Medicamente chimioterapice folosind ASTM D6978.

Rezistența Permeabilității Produselor Chimice	Nivel	Degradare medie (%)
*4% Digluconat de clorhexidină	6	19,0
40% Hidroxid de sodiu (K)	6	-42,9
10-13% Hipoclorit de sodiu	6	14,7
50% Acid sulfuric	6	-20,5
5% Bromură de etidiu	6	3,4
50% Glutaraldehidă	6	27,4
0,1% Fenol	6	33,8
1,5% Soluție apoasă de metanol	6	21,9
3% Iod-Povidonă	6	33,7
10% Percarbonat de sodiu	6	15,4
10% Acid acetic	4	66,7
37% Formaldehidă (T)	3	5,0
30% Peroxid de hidrogen (P)	2	22,8
25% Hidroxid de amoniu (O)	0	-52,0
65% Acid azotic (M)	0	97,6
70% Izopropanol	0	62,2
35% Etanol	0	93,9
99% Acid acetic (N)	0	93,9

* Rata permeabilității 7µg/cm²/min

Nivel de Performanță al Permeabilității și Timpul Măsurat de Străpungere (minute)
0 > * 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indică faptul că mărșă este sub nivelul minim de performanță după cum se indică în EN ISO 374-1:2016 +A1:2018 pentru pericolul dat individual.

50mm

intouch™

POWDER FREE LATEX EXAMINATION GLOVES

EN : USER INFORMATION
DE : NUTZERINFORMATION
FR : FICHE D'INFORMATION
RO : INFORMAȚII UTILIZATOR



Available Size

XS
S
M
L
XL



Last update in August 2022

110mm

EN POWDER FREE LATEX EXAMINATION GLOVES

• Non sterile • Single Use
• Recommended for medical use

Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III) • Medical Devices Regulations 2002 (SI 2002 No. 618, as amended) (UK MDR 2002) • EU Notified Body for (Module B) and Module C2: SATRA Technology Europe Limited (2777) • UK Approved Body for (Module B) and Module C2: SATRA Technology Center Limited (A80321) • EU Declaration of Conformity & UK Declaration of Conformity is accessible at www.intouchcares.com • This information does not reflect the actual duration of protection in the workplace and the differentiation between mixtures and pure chemicals. • The chemical resistance has been assessed under laboratory conditions from samples taken from the palm only (except in cases where the glove is equal to or over 400mm – where the cuff is tested also) and relates only to the chemical tested. It can be different if the chemical used is a mixture. • It is recommended to check that the gloves are suitable for the intended use because the conditions at the workplace may differ from the type test depending on temperature, abrasion and degradation. • When used, protective glove may provide less resistance to the dangerous chemical due to changes in physical properties. Movements, snagging, rubbing, degradation caused by chemical contact etc. may reduce the actual use time significantly. For corrosive chemicals, degradation can be the most important factor to consider in selection of chemical resistant gloves. • Before usage, inspect the gloves for any defect or imperfections. • EN ISO 374-4:2019 Degradation levels indicate the change in puncture resistance of the gloves after exposure to the challenge chemical. • The penetration resistance has been assessed under laboratory conditions and relates only to the tested specimens. • Donning – Hold glove by the bead with one hand. Align the glove thumb with your other hand thumb and slide your hand into the glove, one finger into each glove finger. Pull by the glove palm to get a good fit. Don the other glove by the same procedure. • Doffing – Hold glove bead and pull toward the finger until the glove come off. • Where relevant, a list of the substances contained in the glove which are known to cause allergies, per listed in Annex G of EN ISO 374-5:2016, shall be supplied on request. • Components used in glove manufacturing may cause allergic reactions in some users. If allergic reactions occur, seek for medical advice immediately. • In storage, avoid excessive heat. Open box should be shielded from exposure to direct sun or fluorescent lighting. Gloves are packed in dispenser which is suitable for transport. Keep the gloves in box when not in use.

Resistance to Permeation by Chemicals	Level	Mean Degradation (%)
Diethylamine (G)	0	7.2
96% Sulphuric Acid (L)	0	100.0
40% Sodium Hydroxide (K)	6	-14.9
30% Hydrogen Peroxide (P)	2	-15.6
37% Formaldehyde (T)	1	-22.4

Permeation Performance Level & Measured Breakthrough Time (minutes) 0 > *, 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indicates that the glove falls below the minimum performance level as stated in EN ISO 374-1:2016 +A1:2018 for the given individual hazard.

50mm

FR GANTS D'EXAMEN NITRILES SANS POUVRE

• Non stérile • Usage unique
• Recommandés pour un usage médical

Règlement sur les dispositifs médicaux (UE) 2017/745 et règlement sur les équipements de protection individuelle (UE) 2016/425 (catégorie III). • Règlement sur les dispositifs médicaux de 2002 (SI) 2002 No. 618, tel que modifié (UK MDR 2002) • Organisme notifié de l'UE pour le (module B) et le module C2 est SATRA Technology Europe Limited (2777) • Organisme agréé au Royaume-Uni pour le (module B) et le module C2 est SATRA Technology Center Limited (A80321). • La déclaration de conformité de l'UE et la déclaration de conformité du Royaume-Uni sont accessibles à l'adresse www.intouchcares.com. • Ces informations ne reflètent pas la durée réelle de la protection sur le lieu de travail et la différenciation entre les mélanges et les produits chimiques purs. • La résistance chimique a été évaluée dans des conditions de laboratoire à partir d'échantillons prélevés sur la paume uniquement (sauf dans les cas où le gant mesure 400 mm ou plus – ou la manchette est également testée) et ne concerne que le produit chimique testé. Elle peut être différente si le produit chimique est utilisé dans un mélange. • Il est recommandé de vérifier que les gants sont adaptés à l'utilisation prévue car les conditions sur le lieu de travail peuvent différer de l'état de type en fonction de la température, de l'abrasion et de la dégradation. • Lorsqu'il est utilisé, le gant de protection peut offrir une moindre résistance au produit chimique dangereux en raison des modifications de ses propriétés physiques. Les mouvements, les accrochages, les frottements, la dégradation causée par le contact chimique, etc. peuvent réduire considérablement la durée d'utilisation réelle. Pour les produits chimiques corrosifs, la dégradation peut être le facteur le plus important à prendre en compte dans le choix de gants résistants aux produits chimiques. • Avant l'utilisation, inspectez les gants pour détecter tout défaut ou imperfection. • EN ISO 374-4:2019 Les niveaux de dégradation indiquent le changement de la résistance à la perforation des gants après exposition au produit chimique de référence. • La résistance à la pénétration a été évaluée dans des conditions de laboratoire et ne concerne que les spécimens testés. • Enfilage - Tenez le gant par le talon d'une main. Alignez le pouce du gant avec le pouce de l'autre main et glissez votre main dans le gant, un doigt dans chaque doigt du gant. Tirez sur la paume du gant pour obtenir un bon ajustement. Enfilez l'autre gant en suivant la même procédure. • Enlever le gant - Tenez le talon du gant et tirez vers le doigt jusqu'à ce que le gant se détache. • Le cas échéant, une liste des substances contenues dans le gant et connues pour provoquer des allergies, telles qu'énumérées à l'annexe G de la norme EN ISO 374-5:2016, doit être fournie sur demande. • Les composants utilisés dans la fabrication des gants peuvent provoquer des réactions allergiques chez certains utilisateurs. En cas de réaction allergique, consultez immédiatement un médecin. • Lors du stockage, évitez toute chaleur excessive. La boîte ouverte doit être protégée de l'exposition directe au soleil ou à un éclairage fluorescent. Les gants sont emballés dans un distributeur qui convient au transport. Conservez les gants dans leur boîte lorsqu'ils ne sont pas utilisés.

Résistance à la perméation par les produits chimiques	Niveau	Moyenne Dégradation (%)
Diethylamine (G)	0	7.2
96% Acide sulfurique (L)	0	100.0
40% Hydroxyde de sodium (K)	6	-14.9
30% Peroxyde d'hydrogène (P)	2	-15.6
37% Formaldéhyde (T)	1	-22.4

Niveau de performance de perméation et temps de percée mesurés (minutes) 0 > *, 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indique que le gant est inférieur au niveau de performance minimum tel qu'indiqué dans la norme EN ISO 374-1:2016 +A1:2018 pour le risque individuel donné.

110mm

RO MÂNSUȘI NITRIL NEPUDRATE PENTRU EXAMINARE

• Nesterile • De unică folosință
• Recomandate a fi folosite în scop medical

Regulamentul privind Dispozitivele Medicale (UE) 2017/745 și Regulamentul privind Echipamentul Individual de Protecție (UE) 2016/425 (Categorie III) • Regulamentul privind Dispozitivele Medicale 2002 (SI) 2002 Nr. 618, cu modificări (UK MDR 2002) • Organism Notificat UE pentru (Modulul B) și Modulul C2: SATRA Technology Europe Limited (2777) • Organismul Notificat MB pentru (Modulul B) și Modulul C2: SATRA Technology Center Limited (A80321) • Declarația de Conformitate UE și Declarația de Conformitate MB este accesibilă la adresa www.intouchcares.com • Informația dată nu se referă la protecția locului de muncă și siguranța la diferite amestecuri din produse chimice. • Rezistența chimică a fost evaluată în condiții de laborator pe eșantioane prelevate doar din palmă, cu excepția cazurilor când măsura are 400 mm și mai mult – unde și manșeta este testată) și se referă doar la substanța chimică testată. Rezultatele pot fi diferite dacă este folosit un amestec de substanțe chimice. • Se recomandă de a verifica dacă măsurașile sunt potrivite pentru a fi folosite în scopul propus, deoarece condițiile locului de muncă pot fi diferite de tipul testărilor efectuate și se pot diferenția prin temperatură, abraziune și degradare. • Măsurașile de protecție, în timpul utilizării, pot oferi o rezistență mai mică la substanțele chimice periculoase din cauza modificărilor proprietăților fizice la acțiunea acestor compuși chimici. Mișcarea, strângerea, frecarea, degradarea poate fi cel mai important factor pentru a fi luat în considerare la selecția măsurașilor rezistenți la produsele chimice. • Înainte de folosire, verificați măsurașile la lipsa defectelor și imperfecțiunilor. • EN ISO 374-4:2019 Nivelurile de degradare indică modificarea rezistenței măsurașilor la perforare după expunerea la substanțele chimice. • Rezistența la penetrare a fost evaluată în baza condițiilor de laborator și se referă doar la specișimenele testate. • Înălțarea - Apucați măsurașile de margine cu o mână. Introduceți degetul mare al măsurașii pe degetul mare al celeilalte mâni și trageți mâna în măsuraș, fiecare deget al mâinii în degetul măsurașii. Trageți astfel măsurașul încât să se potrivească. Înălțarea celeilalte măsurașii după același scenariu. • Dezbrăcarea - Țineți marginea măsurașii și trageți de pe degete până măsurașul va ieși. • Unde e cazul, se va oferi la cerere lista componentelor măsurașii care sunt recunoscute ca alergeni, menționate în Anexa G a EN ISO 374-5:2016. • Componentele folosite la producerea măsurașilor pot cauza reacții alergice la unii utilizatori. Dacă se produce o reacție alergică, adresați-vă urgent medicului. • Măsurașile se vor păstra la loc ferit de căldură în exces. Cutia deschisă se va proteja de expunerea directă la soare sau lumină fluorescentă. • Măsurașii sunt ambalați în cutie care se potrivește pentru transportare. Păstrați măsurașile în cutie atunci când nu sunt folosite.

Rezistența Permeabilității Produselor Chimice	Nivel	Degradare medie (%)
Diethylamină (G)	0	7.2
96% Acid sulfuric (L)	0	100.0
40% Hidroxid de sodiu (K)	6	-14.9
30% Peroxid de hidrogen (P)	2	-15.6
37% Formaldehidă (T)	1	-22.4

Nivel de Performanță al Permeabilității și Timpul Măsurat de Strângere (minute) 0 > *, 1 > 10 min, 2 > 30 min, 3 > 60 min, 4 > 120 min, 5 > 240 min, 6 > 480 min

*Indică faptul că măsurașul este sub nivelul minim de performanță, după cum se indică în EN ISO 374-1:2016 +A1:2018 pentru pericolul dat individual.

Comfort and higher dexterity • Chemical permeation tested • Polymer coated • Made with natural rubber latex

Powder-free Examination Gloves

Comfort-care

LATEX

XS

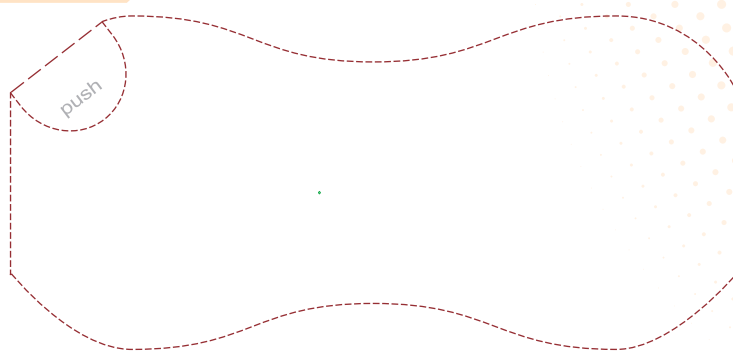
5 - 6
100 gloves
by weight

LATEX

Comfort-care

Powder-free Examination Gloves

PS54



NATURAL COLOUR
XS
5 - 6
100 gloves
by weight

LATEX
Comfort-care
Powder-free Examination Gloves

NATURAL COLOUR
XS
5 - 6
100 gloves
by weight

XS
5 - 6
100 gloves
by weight

NATURAL COLOUR

LATEX
Comfort-care
Powder-free Examination Gloves

XS

LATEX

Comfort-care

Powder-free Examination Gloves

- Gants d'examen en latex sans poudre
- Guderfrei untersuchungshandschuhe aus latex
- Mănuși latex nepudrate pentru examinare

XS

5 - 6
100 gloves
by weight

XS

LATEX

Comfort-care

Powder-free Examination Gloves

In compliance with Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III). Notified Body responsible for PPE EU Type Examination (Module B) and Module C2 On-going Conformity Assessment is SATRA Technology Europe Limited (2777) Bracetown Business park, Clonee, Dublin 15, D15 YN2P, Ireland. EU Declaration of Conformity is accessible at www.intouchcares.com. Product reference: PF NR

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This product contains natural rubber latex which may cause allergic reactions, including anaphylactic responses. This glove product is intended to be worn on the hand to provide a barrier protection against risks associated with contact with certain chemicals, microorganisms & other contaminants.

User information sheet is enclosed in this package.

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UKRP ADVENA LIMITED UK Email info@advenamedical.com
Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

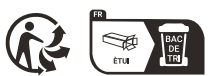
Made in Malaysia



955256614668
REF
IMPS54YWA-XS

Chemicals	Level	Mean degradation
Diethylamine (G)	0	-7.2%
96% Sulphuric acid (L)	0	100.0%
40% Sodium hydroxide (K)	6	-14.9%
30% Hydrogen peroxide (P)	2	-15.6%
37% Formaldehyde (T)	1	-22.4%

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018.
Level 1 represents > 10 min Level 2 represents > 60 min Level 3 represents > 240 min
Level 4 represents > 30 min Level 5 represents > 120 min Level 6 represents > 480 min



Comfort and higher dexterity • Chemical permeation tested • Polymer coated • Made with natural rubber latex

Powder-free Examination Gloves

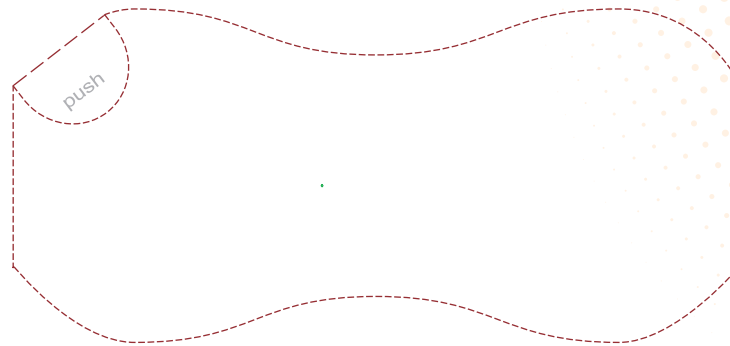
LATEX
Comfort-care

S
6-7
100 gloves
by weight

LATEX
Comfort-care

PS54

Powder-free Examination Gloves



Recommended for medical use



S
6-7
100 gloves
by weight

LATEX
Comfort-care
Powder-free Examination Gloves

NATURAL COLOUR
S
6-7
100 gloves
by weight

S
6-7
100 gloves
by weight

NATURAL COLOUR

LATEX
Comfort-care
Powder-free Examination Gloves

intouch
TM

S

LATEX
Comfort-care

Powder-free Examination Gloves

- Gants d'examen en latex sans poudre
- Guderfrei untersuchungshandschuhe aus latex
- Mănuși latex nepudrate pentru examinare

S
6-7
100 gloves
by weight

S

LATEX
Comfort-care

Powder-free Examination Gloves

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Made in Malaysia

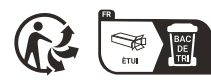


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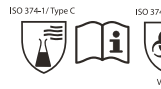
REF
IMPS54YWA-S

Chemicals	Level	Mean degradation
Diethylamine (G)	0	7.2%
96% Sulphuric acid (L)	0	100.0%
40% Sodium hydroxide (K)	6	-14.9%
30% Hydrogen peroxide (P)	2	-15.6%
37% Formaldehyde (T)	1	-22.4%

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min



REF
IMPS54YWA-S



LOT



Comfort and higher dexterity • Chemical permeation tested • Polymer coated • Made with natural rubber latex

Powder-free Examination Gloves

LATEX

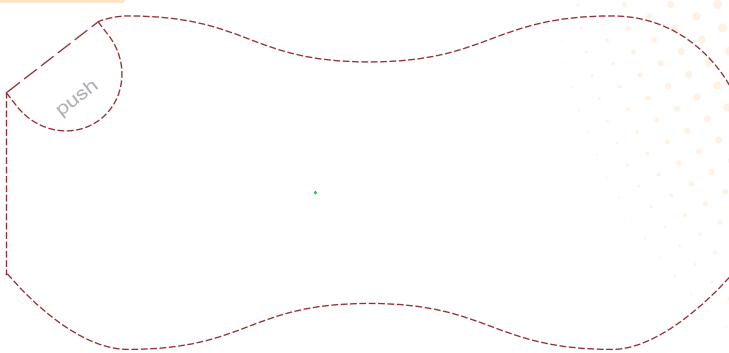
M
7 - 8
100 gloves
by weight

LATEX

Comfort-care

PS54

Powder-free Examination Gloves



M
7 - 8
100 gloves
by weight

LATEX
Comfort-care
Powder-free Examination Gloves

NATURAL COLOUR
M
7 - 8
100 gloves
by weight

M
7 - 8
100 gloves
by weight
NATURAL COLOUR

LATEX
Comfort-care
Powder-free Examination Gloves

intouchTM

M

LATEX

Comfort-care

Powder-free Examination Gloves

- Gants d'examen en latex sans poudre
- Guderfrei untersuchungshandschuhe aus latex
- Mănuși latex nepudrate pentru examinare

M
7 - 8
100 gloves
by weight

M

LATEX

Comfort-care

Powder-free Examination Gloves

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User information sheet is enclosed in this package.

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Pure Offices, Plato Close, Tachbrook Park, Warwick CV34 6WE, United Kingdom.

Made in Malaysia

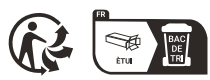


9555256614682

REF
IMPS54YWA-M

Chemicals	Level	Mean degradation
Diethylamine (G)	0	-7.2%
96% Sulphuric acid (L)	0	100.0%
40% Sodium hydroxide (K)	6	-14.9%
30% Hydrogen peroxide (P)	2	-15.6%
37% Formaldehyde (T)	1	-22.4%

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 «A1: 2018.
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min



Comfort and higher dexterity • Chemical permeation tested • Polymer coated • Made with natural rubber latex

Powder-free Examination Gloves

Comfort-care

LATEX

8-9
100 gloves
by weight

LATEX

Comfort-care

Powder-free Examination Gloves

PS54

LATEX
Comfort-care
Powder-free Examination Gloves

NATURAL COLOUR
8-9
100 gloves
by weight



NATURAL COLOUR
8-9
100 gloves
by weight

NATURAL COLOUR
8-9
100 gloves
by weight

LATEX
Comfort-care
Powder-free Examination Gloves

INTOUCH™

LATEX

Comfort-care

Powder-free Examination Gloves

- Gants d'examen en latex sans poudre
- Guderfrei untersuchungshandschuhe aus latex
- Mănuși latex nepudrate pentru examinare

8-9
100 gloves
by weight

LATEX

Comfort-care

Powder-free Examination Gloves

In compliance with Medical Device Regulation (EU) 2017/745 & Personal Protective Equipment Regulation (EU) 2016/425 (Category III). Notified Body responsible for PPE EU Type Examination (Module B) and Module C2. On-going Conformity Assessment is SATRA Technology Europe Limited (2777) Bracetown Business park, Clonee, Dublin 15, D15 YN2P, Ireland. EU Declaration of Conformity is accessible at www.intouchcares.com. Product reference: PF NR

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Made in Malaysia



9555256614699

REF
IMPS54YWA-L

Chemicals	Level	Mean degradation
Diethylamine (G)	0	7.2%
96% Sulphuric acid (L)	0	100.0%
40% Sodium hydroxide (K)	6	-14.9%
30% Hydrogen peroxide (P)	2	-15.6%
37% Formaldehyde (T)	1	-22.4%

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018
Level 1 represents > 10 min Level 3 represents > 60 min Level 5 represents > 240 min
Level 2 represents > 30 min Level 4 represents > 120 min Level 6 represents > 480 min



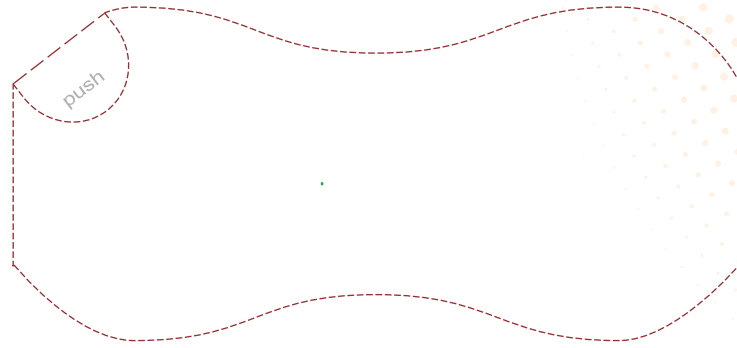
LOT



Comfort and higher dexterity • Chemical permeation tested • Polymer coated • Made with natural rubber latex
 Powder-free Examination Gloves
LATEX
 Comfort-care

XL
 9 - 10
 100 gloves
 by weight

LATEX **PS54**
 Comfort-care
 Powder-free Examination Gloves



LATEX
 Comfort-care
 Powder-free Examination Gloves

NATURAL COLOUR
XL
 9 - 10
 100 gloves
 by weight

XL
 9 - 10
 100 gloves
 by weight

LATEX
 Comfort-care
 Powder-free Examination Gloves

intouchTM

XL

LATEX
 Comfort-care
 Powder-free Examination Gloves

- Gants d'examen en latex sans poudre
- Guderfrei untersuchungshandschuhe aus latex
- Mănuși latex nepudrate pentru examinare

XL
 9 - 10
 100 gloves
 by weight

XL

LATEX
 Comfort-care
 Powder-free Examination Gloves

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Made in Malaysia



9555256614705

REF
 IMPS54YWA-XL

Chemicals	Level	Mean degradation
Diethylamine (G)	0	7.2%
96% Sulphuric acid (L)	0	100.0%
40% Sodium hydroxide (K)	6	-14.9%
30% Hydrogen peroxide (P)	2	-15.6%
37% Formaldehyde (T)	1	-22.4%

Level 0 represents below minimum permeation performance level as stated in EN ISO 374-1: 2016 +A1: 2018
 Level 1 represents > 10 min
 Level 2 represents > 30 min
 Level 3 represents > 60 min
 Level 4 represents > 120 min
 Level 5 represents > 240 min
 Level 6 represents > 480 min

