

Sysmex Europe GmbH · Bornbarch 1 · 22848 Norderstedt · Germany

To whom it may concern

Sysmex Europe GmbH
Bornbarch 1
22848 Norderstedt, Germany
Phone +49 40 527 26-0
Fax +49 40 527 26-100
Info@sysmex-europe.com

### LETTER OF AUTHORIZATION

Whereas Sysmex Europe GmbH ("Sysmex"), who are established, reputable and authorised representative in Europe, Africa and Middle East (EMEA region), officially announced by the manufacturer Sysmex Corporation, having its principal place of business at 1-5-1 Wakinohama-Kaigandoori, Chuo-ku, Kobe 651-0073, Japan, and having the power to grant authorizations to local representatives within the above mentioned markets,

do hereby declare that the company

ECHIPAMED Plus SRL
Valea Trandafirilor 24 "B", off. 80
MD-2001 Chisinau, Moldova (the "COMPANY")

is our distributor and local representative for the following Sysmex products:

Sysmex Haematology- and Urine- Analysers
with Reagents, Accessories, Software and Spare Parts
(the "Products")

In the territory of Moldova (the "TERRITORY")

The **COMPANY** is therefore authorized to carry out all commercial and support activities for the **PRODUCTS** including sales, marketing, application, registration and field service support in the **TERRITORY**.

The **COMPANY** is aware that this special authorisation is limited to the above listed **PRODUCTS** and does not create any further rights for the **COMPANY**.



Company Location Norderstedt Registered AG Kiel HRB 4179 VAT-ID DE 118 687 842 WEEE/ElektroG Reg. Nr. DE 159 56 453 Managing Directors Alain Baverel Seido Biwa Alberto Bonacini Kensuke lizuka Iwane Matsui Stefanie Schaal Jan Willem Schipper Matthias Völkel COMMERZBANK AG, Hamburg IBAN DE 20 2004 0000 0287 1879 00 SWIFT/BIC Code COBADEFFXXX

www.sysmex-europe.com



We hereby grant our warranty following our general conditions of sale for the **PRODUCTS** delivered, consisting of and limited to:

Free of charge supply of spare parts to the **COMPANY** as replacement for defective new parts for a period of 14 months after B/L - AWB date.

This declaration is valid until 31.03.2021 and may be revoked unilaterally by Sysmex in writing before that date for due cause. Such due cause shall, among others, be the termination or expiration of the distributorship relationship, if any, between Sysmex and the **COMPANY**.

On behalf of Sysmex Europe GmbH

Date: March 20th, 2020

Place: 22848 Norderstedt, Germany

Jan-Willern Schipper Senior Executive Officer sysmex

Sysmex Europe GmbH Bornbarch 1 22848 Norderstadt





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To whom it may concern

Sysmex Europe GmbH
Bornbarch 1
22848 Norderstedt, Germany
Phone +49 40 52726-0
Fax +49 40 52726-100
info@sysmex-europe.com

### **DECLARATION**

We, Sysmex Europe GmbH, located at Bornbarch 1, 22848 Norderstedt, Germany, who are established, reputable and authorised representative in Europe (EC REP), Africa and Middle East (EMEA region), officially announced by the manufacturer Sysmex Corporation, Japan hereby confirm that our Haematology Analysers

#### XN-1000, XN-2000, XP-300 and UX-2000

are 'closed systems' and only to be used together with Sysmex Reagents, Sysmex Controls and Sysmex Calibrators. Every change of this closed system by the user is regarded as 'non-specified use' by Sysmex.

The technology of all Sysmex IVD analysers is fine-tuned together with the corresponding reagents used on each single analyser. Thereby, using Sysmex reagents maintains optimum performance as well as optimal and enhanced accuracy of the system. There is a high interdependency between research and using/finding optimal reagents for any new parameter(s). As Sysmex is actively doing research, it is thereby ensured that Sysmex reagents fulfil best practice requirements for any research parameter(s), which later will become diagnostic parameter(s) after the legally required procedures under Annex VIII-IVD-Directive 98/79/EC.

Therefore Sysmex Reagents offer best performance on Sysmex Analysers.

Sysmex Europe GmbH

The Reagents, Controls and Calibrators listed on the following page are allowed to be used on Sysmex Haematology Analysers.

On behalf of Sysmex Europe God

Date: January 14th, 2016

Place: 22848 Norderstedt, Germany

i.A. Katharina Paucke

Manager Regulatory Affairs

"Design and specifications may be subject to changes due to further product development. Changes are confirmed by their appearance on a newer document and verification according to its date of issue."





Reagents, Controls and Calibrators that are allowed to be used on Sysmex Haematology Analysers:

XN-1000	XN-2000	XP-300	UX-2000
CELLPACK DCL	CELLPACK DCL	CELLPACK	UX II PACK-BAC
CELLPACK DST	CELLPACK DST	STROMATOLYSER-WH	UX II PACK-SED
CELLPACK DFL	CELLPACK DFL	CELLCLEAN	UX II SEARCH -BAC
Lysercell WDF	Lysercell WDF	EIGHTCHECK-3WP	UX II SEARCH -SED
Lysercell WNR	Lysercell WNR	SCS-1000	UX II SHEATH
Lysercell WPC	Lysercell WPC		UX CLEAN -C
SULFOLYSER	SULFOLYSER	A STATE OF THE PARTY OF THE PAR	MEDITAPE II 10U
Fluorocell PLT	Fluorocell PLT		MEDITAPE II 9U
Fluorocell RET	Fluorocell RET	TO A RESERVED BY	MEDITAPE II 10K
Fluorocell WDF	Fluorocell WDF		UF II CONTROL
Fluorocell WNR	Fluorocell WNR		MEDITAPE CHECK 1
Fluorocell WPC	Fluorocell WPC		MEDITAPE CHECK 2
CELLCLEAN	CELLCLEAN	<b>等于各种区的</b> 设备等等的的	UF II Calibrator
CELLCLEAN AUTO	CELLCLEAN AUTO	Public and the second second	Transportation 2
XN CHECK	XN CHECK		
XN CHECK BF	XN CHECK BF	<b>一生,其一种</b>	10000000000000000000000000000000000000
XN CAL	XN CAL		
XN CAL PF	XN CAL PF	In the Country of the Assessment	

End of list





Sysmex Europe GmbH · Bornbarch 1 · 22848 Norderstedt · Germany

To whom it may concern

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## **Composition of Sysmex Reagents**

The composition of Sysmex Reagents is highly confidential! Therefore only active components and those classified as dangerous must be declared on the product labelling.

The below listed table gives an overview of these components in Sysmex Reagents:

Cellpack	Sodium chloride 6.38 g/L Boric acid 1.0 g/L Sodium tetraborate 0.2 g/L EDTA-2K 0.2 g/L				
CELLPACK DCL	Sodium chloride 0.7% Tris buffer 0.2% EDTA-2K 0.02%				
CELLPACK DST	Sodium chloride 15.7% Tris buffer 4.3% EDTA-2K 0.4%				
CELLPACK DFL	Tricine buffer 0.17%				
CELLCLEAN	Sodium Hypochlorite (available chlorine concentration 5.0%)				
CELLCLEAN AUTO	Sodium Hypochlorite (available chlorine concentration 5.0%)				
Stromatolyser-WH	Organic quaternary ammonium salt 8.5 g/L Sodium chloride 0.6 g/L				
Lysercell WDF	Organic quaternary ammonium salts 0.07% Nonionic surfactant 0.17%				
Lysercell WNR	Organic quaternary ammonium salts 0.20% Nonlonic surfactant 0.10%				
Lysercell WPC	Anionic surfactant 0.03% Nonionic surfactant 0.12%				
Sulfolyser	Sodium lauryl sulfate 1.7 g/L				
Fluorocell PLT	Oxazine 0.003% Ethylene glycol 99.9%				
Fluorocell RET	Polymethine 0.03% Methanol 7.9% Ethylene glycol 92.0%				
Fluorocell WDF	Polymethine 0.002% Methanol 3.0% Ethylene glycol 96.9%				
Fluorocell WNR	Polymethine 0.005% Ethylene glycol 99.9%				



Company Location Norderstedt Registered AG Kiel HRB 4179 VAT-ID DE 118 687 842 WEEE/ElektroG Reg. Nr. DE 159 56 453 Managing Directors Fernando Andreu Kensuke lizuka Takeshi Kubota Kazuya Obe Dr. Michael Schaefer Dr. Jürgen Schulze Matthias Völkel The Bank of Tokyo-Mitsubishi UFJ, Ltd. Hamburg Bank ID-Code 300 107 00 Account Nr. 03 77 13 IBAN DE03 3001 0700 0000 0377 13 SWIFT/BIC Code BOTKDEDX





Fluorocell WPC	Polymethine 0.004% Ethanol 15.1%						
	Ethylene glycol 84.8%						
XN CHECK	quality control material; includes stabilized human red blood cells, human white blood cells, a platelet and nucleated red blood cell component in a preservative medium.						
XN CHECK BF	quality control material; includes stabilized human red blood cells and white blood cells in a preservative medium.						
XN CAL	calibrator; includes stabilized human red blood cells, human white blood cells, a platelet and nucleated red blood cell component in a preservative medium.						
XN CAL PF	calibrator; includes stabilized human red blood cells and a platelet component in a preservative medium.						
Eightcheck-3WP	quality control material; includes stabilized human red blood cells, fixed mammalian white blood cells and a platelet component in a preservative medium						
SCS-1000	quality control material; contains stabilised human red blood cells, fixed mammalian white bloodcells, and a platelet component in a medium containing preservatives.						
UX II PACK-BAC	Buffer 1.9% Cation surfactant 0.1%						
UX II PACK-SED	Buffer 2.1%						
UX II SEARCH -BAC	Polymethine Dye 0.01% ( w / w ) Ethylene glycol 99.9% ( w / w )						
UX II SEARCH -SED	Polymethine Dye 0.03% ( w / w ) Ethylene glycol 99.9% ( w / w )						
UX II SHEATH	Tris Buffer 0.14%						
UX CLEAN -C	t-Octylphenoxypolyethoxyethanol < 1.0 %  Sodium azide < 0.1 %  Sodium phosphoto tribasia dedocabydrata < 1.0 %						
	Sodium phosphate tribasic dodecahydrate < 1.0 %  Reactive ingredients (per 100 test strips)						
	[Glucose] Glucose oxidase: 700 I.U., Peroxidase: 175 P.U., 4-Aminoantipyrine: 14.0 mg, 1-Naphthol-3,6-disufonic acid, disodium salt: 14 mg						
	[Protein] Tetrabromophenol blue: 0.35 mg						
	[Bilirubin] 2-Methyl-5-nitroaniline: 1.9 mg, Sodium nitrite: 1.0 mg						
	[Urobilinogen] 3,3'-Dimethoxy-4,4'-biphenylbis (diazonium tetrafluoroborate): 0.16 mg						
MEDITAPE II 10U	[Creatinine] 2,6-Dichloro-4'-hydroxy-3',3"-dimethyl-3-sulfofuchsone-5',5"-dicarboxylic acid, trisodium salt: 0.34 mg, Palladium (II) chloride: 0.10 mg						
	[pH] Bromocresol green: 0.07 mg, Bromoxylenol blue: 0.72 mg						
	[Blood] Cumene hydroperoxide: 30.0 mg, 3,3',5,5'-Tetramethylbenzidine: 15.0 mg						
	[Ketones] Sodium nitroprusside: 12.0 mg						
	[Nitrite] Sulfanilamide: 3.9 mg, N-1-Naphthylethylenediamine dihydrochloride: 0.3 mg						
	[Leukocytes] 3-(N-Toluenesulfonyl-L-alanyloxy) indole: 0.69 mg, 2-Methoxy-4-(N-morpholino)benzenediazonium: 0.38 mg						





	Reactive ingredients (per 100 test strip	os)				
	[Glucose] Glucose oxidase: 700 I.U., Per 4-Aminoantipyrine: 14.0 mg, 1-Naphthol-					
	[Protein] Tetrabromophenol blue: 0.35 m	g				
	[Bilirubin] 2-Methyl-5-nitroaniline: 1.9 mg	, Sodium nitrite: 1.0 mg				
MEDITAPE II 9U	[Urobilinogen] 3,3'-Dimethoxy-4,4'-bipher	nylbis (diazonium tetrafluoroborate): 0.16 mg				
	[pH] Bromocresol green: 0.07 mg, Bromo	exylenol blue: 0.72 mg				
	[Blood] Cumene hydroperoxide: 30.0 mg	, 3,3',5,5'-Tetramethylbenzidine: 15.0 mg				
	[Ketones] Sodium nitroprusside: 12.0 mg					
	[Nitrite] Sulfanilamide: 3.9 mg, N-1-Naphi	thylethylenediamine dihydrochloride: 0.3 mg				
	[Leukocytes] 3-(N-Toluenesulfonyl-L-alar morpholino)benzenediazonium: 0.38 mg	nyloxy) indole: 0.69 mg, 2-Methoxy-4-(N-				
	Reactive ingredients (per 100 test strip	M = Q :				
	[Glucose] Glucose oxidase: 700 I.U., Per 4-Aminoantipyrine: 14.0 mg	oxidase: 175 P.U.,				
	[Protein] Tetrabromophenol blue: 0.35 mg	[Protein] Tetrabromophenol blue: 0.35 mg				
	[Albumin] 4,5,6,7-Tetrachloro-2',4',5',7'-te	[Albumin] 4,5,6,7-Tetrachloro-2',4',5',7'-tetraiodoflurescein disodium salt: 0.14 mg				
	[Bilirubin] 2-Methyl-5-nitroaniline: 1.9 mg, Sodium nitrite: 1.0 mg					
MEDITAPE II 10K	[Creatinine] 2,6-Dichloro-4'-hydroxy-3',3"-dimethyl-3-sulfofuchsone-5',5"-dicarboxylic acid, trisodium salt: 0.34 mg, Palladium (II) chloride: 0.10 mg					
	[pH] Bromocresol green: 0.07 mg, Bromoxylenol blue: 0.72 mg					
	[Blood] Cumene hydroperoxide: 30.0 mg, 3,3',5,5'-Tetramethylbenzidine: 15.0 mg					
	[Ketones] Sodium nitroprusside: 12.0 mg	[Ketones] Sodium nitroprusside: 12.0 mg				
	[Nitrite] Sulfanilamide: 3.9 mg, N-1-Naphthylethylenediamine dihydrochloride: 0.3 mg					
	[Leukocytes] 3-(N-Toluenesulfonyl-L-alar morpholino)benzenediazonium: 0.38 mg	[Leukocytes] 3-(N-Toluenesulfonyl-L-alanyloxy) indole: 0.69 mg, 2-Methoxy-4-(N-morpholino)benzenediazonium: 0.38 mg				
	UF II CONTROL -H	UF II CONTROL -L				
UF II CONTROL	Control particles 0.4% ( w / w ) NOTE: This product contain Latex particle.	Control particles 0.1% ( w / w ) NOTE: This product contain Latex particle.				
	prepared from human urine;	prepared from human urine;				
MEDITAPE CHECK 1	Chemical and biochemical substances as well as constituents of human origin are contained.					
	prepared from human urine;	prepared from human urine;				
MEDITAPE CHECK 2	Chemical and biochemical substances as well as constituents of human origin are contained.					
Control particles 0.4% ( w / w )						
UF II Calibrator	NOTE : This product contain Latex partic	le.				

On behalf of Sysmex Europe GmbH

i.A. Katharina Paucke

Manager Regulatory Affairs

Date: January 19th, 2016

Place: 22848 Norderstedt, Germany

Sysmex Europe GmbH Bornbarch 1 22848 Norderstedt

"Design and specifications may be subject to changes due to further product development. Changes are confirmed by the reached on a newer document and verification according to its date of issue."



## Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

SYSMEX EUROPE GmbH Bornbarch 1 22848 Norderstedt Deutschland

has established and applies a quality management system for medical devices for the following scope:

see attachment

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance.

Effective Date:

2019-05-17

Certificate Registration No.:

SX 60137613 0001

An audit was performed. Report No.: 21245244 005

This Certificate is valid until:

2022-05-16

Certification Body



Date 2019-04-29

TÜV Rheinland LGA Products GmbH Tillystraße 2/90431 Nürnberg

Tel.: +49 221 806-1371 Fax: +49 221 806-3935 e-mail:cert-val

# Certificate

Standard

ISO 14001:2015

Certificate Registr. No.

01 104 110072

Certificate Holder:

SYSMEX EUROPE GmbH

Bornbarch 1 22848 Norderstedt Germany

Scope:

Sales, marketing and service of in-vitro diagnostic medical devices

Proof has been furnished by means of an audit that the requirements of ISO 14001:2015 are met.

Validity:

The certificate is valid from 2020-07-25 until 2023-07-24.

First certification 2011

2020-03-19

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln

land®



DAKKS

Deutsche
Akkreditierungsstelle
D-ZM-16031-01-00







## Application of Directives:

- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means	of	conf	form	itv.
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The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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- 1			•			16			21		

Product name:

CELLPACK DCL

Classification:

Other device (except Annex II and self-testing devices)

## List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

1	To reference or construction	1 B /	A	-	_ 1	
ı	Lega	I\/	lan	IIII	CIL	irer.

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

Phiroshi Yamane, Executive Vice President

Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Børnbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: HARCH ZUT 2018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until

25.05.2022 or until a revised declaration is Issued due to product modifications.



1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan Tel. +81-78-265-0500 Fax. +81-78-265-0524



## Application of Directives:

- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

### Means of conformity:

The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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	100	luct	IUC	111111	Callo	11.

Product name:

SULFOLYSER

Classification:

Other device (except Annex II and self-testing devices)

## List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

ı	Lega	I N/	lanut	factu	rer

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

Hiroshi Yamane, Executive Vice Preside

#### Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: MARCH 21st 2018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.



## Application of Directives:

- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

## Means of conformity:

The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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Г	100	uci	lue	111111	IUali	OII.

Product name:

Lysercell WNR

Classification:

Other device (except Annex II and self-testing devices)

## List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

## Legal Manufacturer:

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

Date: 13 March, 2018 Hiroshi Yamane, Executive Vice President

#### Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Børnbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: HARCH ZIST 7018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.





## Application of Directives: - 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices Means of conformity: The following product is in conformity with - Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive. Product identification: Product name: Lysercell WDF Classification: Other device (except Annex II and self-testing devices) List of Applied Standards: - Harmonised Standards used for conformity assessment are listed in the technical documentation. Legal Manufacturer: Name: SYSMEX CORPORATION 1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan Address: Hiroshi Yamano, Executive Vice President Authorised officer:

Authorised officer: Date: MARCH 21 ST 2018

Fernando Andreu, Chief Operations Officer

SYSMEX EUROPE ØMBH

Bornbarch 1,22848 Norderstedt, Germany

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

Authorised representative:

Name:

Address:



Application of Directive	es:
- 98/79/EC of 27 Octob	per 1998 on In Vitro Diagnostic Medical Devices
Means of conformity:  The following product - Directive 98/79/EC ba Annex III of the Direct	ased on the conformity assessment procedures in accordance with
Product identification: Product name: Classification:	Fluorocell WNR  Other device (except Annex II and self-testing devices)
List of Applied Standard - Harmonised Standard documentation.	rds: s used for conformity assessment are listed in the technical
Legal Manufacturer: Name: Address:	SYSMEX CORPORATION  1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan
Authorised officer:	Hiroshi Yamane, Executive Vice President
Authorised representat	tive:
Name:	SYSMEX EUROPE GMBH
Address:	Bornbarch 1, 22848 Norderstedt, Germany
Authorised officer:	Fernando Andreu, Chief Operations Officer
This declaration of conformity is 25.05.2022 or until a revised decl	issued under the sole responsibility of the manufacturer and is valid until aration is Issued due to product modifications.



Application	of	<b>Directives</b>
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- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means	of	conf	fo	rm	ity	:
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The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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	100	UOL	I		1100	LIOI	ι.

Product name:

Fluorocell WDF

Classification:

Other device (except Annex II and self-testing devices)

### List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

Legal Manufacturer:

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

Hiroshi Yamane, Executive Vice President

Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: MARCH 21 17 2018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until

25.05.2022 or until a revised declaration is Issued due to product modifications.



## Application of Directives: - 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices Means of conformity: The following product is in conformity with - Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive. Product identification: Product name: CELLPACK DFL Classification: Other device (except Annex II and self-testing devices) List of Applied Standards: - Harmonised Standards used for conformity assessment are listed in the technical documentation. Legal Manufacturer: Name: SYSMEX CORPORATION 1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan Address: Date: 13 March, 2018 Hiroshi Yamane, Executive Vice President

Authorised representative:

Authorised officer:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date:

MARCHZII 2018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.





Application of Directive	es: per 1998 on In Vitro Diagnostic Medical Devices
- 98/19/EC 01 27 Octob	Set 1998 off the Vitto Diagnostic Medical Devices
Means of conformity: The following product - Directive 98/79/EC by Annex III of the Direct	ased on the conformity assessment procedures in accordance with
Product identification: Product name:	Fluorocell RET
Classification:	Other device (except Annex II and self-testing devices)
oradomoution.	
List of Applied Standard - Harmonised Standard documentation.	rds: Is used for conformity assessment are listed in the technical
Legal Manufacturer:	SYSMEX CORPORATION
Address:	1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan
Authorised officer:	Hiroshi Yamane, Executive Vice President
Authorised representa	tive:
Name:	SYSMEX EUROPE GMBH
Address:	Bornbarch 1, 22848 Norderstedt, Germany
Authorised officer:	Date: HARCH ZIT 2018

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

Fernando Andreu, Chief Operations Officer



Application of Directive - 98/79/EC of 27 Octob	es: ber 1998 on In Vitro Diagnostic Medical Devices
Means of conformity:  The following product  Directive 98/79/EC b  Annex III of the Direct	ased on the conformity assessment procedures in accordance with
Product identification: Product name:	CELLCLEAN
Classification:	Other device (except Annex II and self-testing devices)
List of Applied Standard - Harmonised Standard documentation.	rds: Is used for conformity assessment are listed in the technical
Legal Manufacturer: Name: Address:	SYSMEX CORPORATION  1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan  Date: 13 March 2018
Authorised officer:	Hiroshi Yamane, Executive Vice President
Authorised representa	
Name:	SYSMEX EUROPE GMBH
Address:	Bornbarch 1, 22848 Norderstedt, Germany
Authorised officer:	Date: MARCH 2(4 2018  Fernando Andreu Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.



## Application of Directives:

- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

## Means of conformity:

The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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Product name:

XN CHECK

Classification:

Other device (except Annex II and self-testing devices)

### List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

Legal Manufacturer:

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

Tiroshi Yamane, Executive Vice Presiden

Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: HARCH 211 7018

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.





## Application of Directives:

- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

## Means of conformity:

The following product is in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive.

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Product name:

XN CAL

Classification:

Other device (except Annex II and self-testing devices)

### List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

#### Legal Manufacturer:

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan

Authorised officer:

### Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: HARCH 21st 7015

Fernando Andreu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

