

GMED certifie que le système de management de la qualité développé par

GMED certifies that the quality management system developed by

ELITECH CLINICAL SYSTEMS SAS Zone Industrielle 61500 SEES FRANCE

pour les activités

for the activities

Conception, production, contrôle et commercialisation de produits de chimie cliniques pour le diagnostic in vitro. Validation de la combinaison réactifs et automates. Distribution d'automates et de produits de chimie cliniques pour le diagnostic in vitro.

Design, production, control and sales of clinical chemistry products intended to be used for in vitro diagnostics. Validation of the combination reagents and analyzers. Distribution of clinical chemistry analyzers and products for in vitro diagnostics.

réalisées sur le(s) site(s) de performed on the location(s) of

ELITech Clinical Systems SAS Zone industrielle - 61500 SEES - FRA

est conforme aux exigences des normes internationales complies with the requirements of the international standards

NF EN ISO 13485 : 2016

Début de validité / *Effective date :* July 28th, 2020 (included) Valable jusqu'au / *Expiry date :* July 27th, 2023 (included) Etabli le / *Issued on :* July 17th, 2020



GMED N° 10462–7 Ce certificat est délivré selon les règles de certification GME

Ce certificat est délivré selon les règles de certification GMED / This certificate is issued according to the rules of GMED certification

Renouvelle le certificat 10462-6

GMED • Société par Actions Simplifiée au capital de 300 000 € • Organisme Notifié/Notified Body n° 0459 Siège social : 1, rue Gaston Boissier - 75015 Paris • Tél. : 01 40 43 37 00 • gmed.fr



ELITech Clinical Systems Zone industrielle 61500 Sées - France Tél : +33 (0)2 33 81 21 00 Fax : +33 (0)2 22 28 77 51 www.elitechgroup.com



DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone industrielle 61500 SEES France, déclarons sous notre seule responsabilité que les réactifs référencés dans la liste ci-jointe (2 pages), sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.

Ces dispositifs sont classés dans la catégorie « autre dispositif » puisqu'ils n'appartiennent ni à la liste A et liste B de l'annexe II et ni à la classe des autotests.

Cette déclaration est basée sur le contenu de chaque dossier technique et s'appuie sur la certification de notre système qualité selon la norme NF EN ISO 13485 : 2016 (Certification valable jusqu'au 27 juillet 2023).

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents such as listed attached (2 pages), conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to in vitro diagnostic medical devices and to the public health code.

These devices are classified in the "other device" category since they do not belong neither to list A or list B of annex II nor to self-testing class.

This declaration is based on the contents of each technical file and is supported by the certification of our quality system according to the standard NF EN ISO 13485 : 2016 (Certification valid until July 27th , 2023).

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos referenciados en la lista adjunta (2 páginas), son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico in vitro y el código de salud pública.

Estos dispositivos se clasifican en la categoría "otro dispositivo", ya que no pertenecen a la lista A ni a la lista B del anexo II, tampoco a la clase de autodiagnóstico.

Esta declaración se basa en el contenido de cada expediente técnico y está respaldado por la certificación de nuestro sistema de calidad según la norma NF EN ISO 13485 : 2016 (Certificación válida hasta el 27 de Julio 2023).

Sées, le 12 Mai 2021

Valérie LAMBERT, Responsable des Affaires Réglementaires Regulatory Affairs Manager Responsable de los Asuntos Reglementario	ELITech Clinical Systems SAS Zone Industrielle 61500 SEES - France I. : +33(0)2 33 81 21 00 - Fax : +33(0)2 33 28 77 51 SIRET 318 365 228 00036	Cécile GOUBAULT, Directeur Général Délégué Managing Director Directora General
Toutet		

Société par actions simplifiée au capital de 1.688.392,33 € - SIREN : 318 365 228 - RCS ALENCON

REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCES - REFERENCIAS	Code GMDN		
M	etabolites divers / Miscellaneous metabolites			
ALBUMIN	ALBU-0600/0700/0250/MB30	53597		
ALBUMIN ENVOY BILIRUBIN DIRECT 4+1	ALBU-0850			
ILIRUBIN DIRECT 4+1	BIDI-0600/0250 BITO-0600/0250	53233		
ILIRUBIN TOTAL & DIRECT 4+1	BITD-0600	53229		
REATININE ENVOY	CRSL-0850	53250		
REATININE JAFFE	CRCO-0600/0700	53251		
	CRSL-M490	53250		
REATININE PAP SL	CRSL-0630/0250	04-732802.0		
IRECT BILINUBIN ENVOY	BIDI-M430 BIDV-0850	53233		
LUCOSE ENVOY	GPSL-0850	53233		
LUCOSE HK	-1			
LUCOSE HK SL	GHSL-0600/0250	53301		
LUCOSE PAP	GPSL-M690			
LUCOSE PAP SL	GPSL-0507/0500/0707/0700/0250/0455/0497			
ICROPROTEIN PLUS	LACT-0100 PRTU-0600/0250	53342		
HOSPHORUS	PHOS-0600/0230/M430			
HOSPHORUS ENVOY	PHOS-0850	59123		
DTAL BILIRUBIN	BITO-M430	53229		
DTAL BILIRUBIN ENVOY	BITV-0850	53229		
DTAL PROTEIN DTAL PROTEIN ENVOY	PROB-M830	-		
DTAL PROTEIN PLUS	PROB-0850	53985		
REA	PROB-0600/0700/0250 URSL-M830			
REA ENVOY	URSL-0850	53587		
REA UV SL	URSL-0427/0420/0500/0507/0250/0455			
RIC ACID	AUML-M830			
	AUVD-0850	53583		
RIC ACID MONO SL	AUML-0497/0427/0420/0500/0507/0707/0250	00000		
	AUSL-0250			
	PRTU-M230	53481		
	Enzymes / Enzymes			
P (DEA) SL	PASL-0400/0420/0230			
PENVOY	PIVD-0850	52928		
PIFCC	ALPI-0230			
T ENVOY	ALSL-0850 ALSL-M490	50000		
T/GPT 4+1 SL	ALSL-0410/0430/0510/0250/0455	52923		
IYLASE	AMSL-04430			
IYLASE ENVOY	AMSL-0850	52940		
IYLASE SL	AMSL-0390/0400/0230			
T/GOT	ASSL-M490			
T ENVOY T/GOT 4+1 SL	ASVD-0850	52954		
IOLINESTERASE	ASSL-0410/0430/0510/0250/0455 CHES-0053			
ENVOY	CKSL-0850	52971 53003		
-MB ENVOY	CMSL-0850			
-MB SL / CKMB	CMSL-0410/0430/0230	- 52994		
NAC	CKSL-M230	53003		
NAC SL	CKSL-0410/0430/0230	33003		
MMA-GT MMA-GT PLUS SL	GISL-M230	-		
T ENVOY	GISL-0400/0420/0250 GISL-0850	53027		
1 ENVOY	LLSL-0850			
1 IFCC	LLSL-M230	53072		
1-L SL	LLSL-0400/0420/0230			
ASE	LPSL-0250			
ASE ENVOY	LPSL-0850	53108		
	es / Oligo-élements / Electrolytes / Trace-elements			
CIUM ARSENAZO	CALA-0600/0250/M430	45789		
CIUM ENVOY	CALA-0850			
ORIDE NENVOY	CHLO-0600/0250	60037		
N FERENE	FEFE-0330/0500/0220	54758		
SNESIUM ENVOY	EFEF-0230/0600/M230 MAGX-0850			
SNESIUM XB	MGXB-0250/0600/M430	46795		
GNESIUM XYLIDYL	MAGX-0230/0600	-		
	Lipides / Lipids	A CONTRACTOR OF A CONTRACT		
DLESTEROL	CHSL-M690			
LESTEROL ENVOY	CHSL-0850	53359		
LESTEROL HDL SL 2G	HDLL-0230/0380/0390	53391		
LESTEROL LDL SL 2G	LDLL-0230/0380/0390	53395		
LESTEROL SL	CHSL-0507/0500/0700/0707/0250/0455/0497	53359		
CHOLESTEROL	CHDL-0250/0600/M330			
CHOLESTEROL ENVOY	HDLL-0850	53391		
	CLDL-0250/M330	53395		
CHOLESTEROL ENVOY	LDLL-0850	00000		
		TGML-0850 TGML-0427/0425/0515/0700/0517/0707/0497 53460		
SLYCERIDES MONO SL NEW				

Vie

REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCES - REFERENCIAS	Code GMDN		
Contrôles-Ca	librants-Standards / Controls-Calibrators-Standards			
OLESTEROL HDL 2G CALIBRATOR	HDLL-0011/0041	44696		
HOLESTEROL LDL 2G CALIBRATOR	LDLL-0011/0041	41728		
IOLESTEROL Standard 200 mg/dL	CHOL-0055	44698		
(-MB CONTROL	CKMB-0900	44693		
ICAL 2	CALI-0550	47868		
ITROL I	CONT-0060	47869		
ITROL II	CONT-0160			
LUCOSE Standard 100 mg/dL	GLUP-0055	41818		
DL LDL CALIBRATOR	HLCA-0041	47868		
E CONTROL I	ISCT-0046	47869		
E CONTROL II	ISCT-0047	52402		
ICROPROTEIN PLUS Standard 100 mg/dL	PRTU-0022	53482 44702		
RIGLYCERIDES Standard 200 mg/dL	TRIG-0055	53588		
REA Standard 50 mg/dL	URUV-0055	44704		
RIC ACID Standard 6 mg/dL	ACUR-0055	44704		
	Protéines spécifiques / Specific proteins	50055		
ITI-STREPTOLYSIN O	ASLO-0250	59055 53705		
	ICRP-0400/M230 ICRP-0043	41838		
RP IP CALIBRATOR SET	ICRP-0043			
RP IP CONTROL I	ICRP-0046	41839		
RP IP CONTROL II	CRPW-0230	53705		
RP WR	CRPW-0230 CRPW-0043	41838		
RP WR CALIBRATOR SET	CRPW-0043	41839		
RP WR CONTROL	CRPW-0045 CRPW-0850	53705		
	IFRT-0230	53718		
RRITIN CALIBRATOR	IFRT-0042	41927		
APTOGLOBIN IP	IHAP-0400	53737		
bA1c	HBAC-0240	59090		
bAIC CALIBRATOR SET	HBAC-0043	53315		
ATC CONTROL L + H	HBAC-0049	44435 53760 53787 53795 53475 53475		
AIP	IIGA-0400			
GIP	IIGG-0400			
MIP	IIGM-0400			
ALBUMIN IP	IMAL-0400			
ALBUMIN IP CALIBRATOR SET	IMAL-0043			
ALBUMIN IP CONTROL I	IMAL-0046			
ALBUMIN IP CONTROL II	IMAL-0047	53478		
ROSOMUCOID IP	IORO-0400	53606 53957		
REALBUMIN IP	IPAL-0400			
ROTEIN IP CALIBRATOR SET	IPRO-0043	53593		
CALIBRATOR	IRFA-0042	42230		
HEUMATOID FACTOR	IRFA-0230	55111		
HEUMATOLOGY CONTROL I	IRCT-0046	47869		
HEUMATOLOGY CONTROL II	IRCT-0047	47005		
RANSFERRIN IP	ITRF-0400	59041		
	Vitamines/Vitamins			
TAMIN D	VITD-0250	54476		
TAMIN D CALIBRATOR SET	VITD-0043	54474		
TAMIN D CONTROL SET	VITD-0049	54475		
	Solutions pour électrodes selectives d'ions /			
	SE Solutions for ion-selective electrodes	59238		
E BASELINE SOLUTION ENVOY	ISBA-0850			
E CALIBRATORS	ISCA-0250			
E CALIBRATOR ENVOY	ISCV-0850	59058		
E CLEANER/CONDITIONER	ISCC-0280	1 with the second		
E DILUENT	ISDI-0250			
E DILUENT ENVOY	ISDV-0850			
E REFERENCE SOLUTION	ISRS-0800	59238		
E REFERENCE SOLUTION ENVOY	vage pour les équipements ELITech Clinical Systems /			
	olutions for ELITech Clinical Systems Equipments			
		59058		
CID SOLUTION for ELITech Clinical Systems Analyzers	SLHC-5900	59058		
YSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers	SLNA-5900	a tracket to see which have		
YSTEM SOLUTION	SLSY-5905	58236		
YSTEM SOLUTION for ELITech Clinical Systems Analyzers	SLSY-5900 SOLA-M163	59058		
ASH SOLUTION A	WASH SOLUTION B	59058		
ASH SOLUTION B	ests d'agglutination / Agglutination tests			

Vla



Current issue date:
Expiry date:
Certificate identity number

22 June 2021 21 June 2024 10361225 Original approval(s): ISO 13485 - 9 June 2019

Certificate of Approval

This is to certify that the Management System of:



Van Rensselaerweg 4, 6956 AV Spankeren, The Netherlands

has been approved by Lloyd's Register to the following standards:

ISO 13485:2016

Approval number(s): ISO 13485 - 00020722

This certificate is valid only in association with the certificate schedule bearing the same number on which the locations applicable to this approval are listed.

The scope of this approval is applicable to:

Design, development and manufacturing of clinical chemistry analyzers, contract manufacturing of erythrocyte sedimentation rate analyzers and warehousing of erythrocyte sedimentation rate tubes for the in vitro diagnostic investigation of samples of human origin.

Paul Graaf Chief Operating Officer, Management Systems, MSIS Issued by: Lloyd's Register Nederland B.V. for and on behalf of: Lloyd's Register Quality Assurance Limited



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Certificate identity number: 10361225

Certificate Schedule

Location

Activities

ELITechGroup B.V.

ISO 13485:2016

Design, development and manufacturing of clinical chemistry analyzers, contract manufacturing of erythrocyte sedimentation rate analyzers and warehousing of erythrocyte sedimentation rate tubes for the in vitro diagnostic investigation of samples of human origin.

ELITechGroup B.V.

Kanaaldijk 90, 6956 AX Spankeren, The Netherlands

Van Rensselaerweg 4, 6956 AV Spankeren, The Netherlands

ISO 13485:2016

Design, development and manufacturing of clinical chemistry analyzers, contract manufacturing of erythrocyte sedimentation rate analyzers and warehousing of erythrocyte sedimentation rate tubes for the in vitro diagnostic investigation of samples of human origin.



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ELITechGroup B.V. P.O.Box 100 6950 AC Dieren Van Rensselaerweg 4 6956 AV Spankeren The Netherlands T: +31 313 430 500 info.ecsnl@elitechgroup.com www.elitechgroup.com Chamber of Commerce 09175642

Spankeren, 16 April 2021

To Whom It May Concern

MANUFACTURER'S AUTHORIZATION LETTER

We, **ELITechGroup B.V**., manufacturer of automated clinical chemistry analyzers, having factories at: Van Rensselaerweg 4 6956 AV Spankeren The Netherlands

and being a company of the ELITechGroup hereby confirm that:

GBG-MLD SRL

Str. Tighina 65, of. 607 Mun. Chişinău, MD-2001 Moldova

is our distributor in Moldova and is fully authorized to offer and deliver the ELITechGroup B.V. products as mentioned in Appendix A.

GBG-MLD SRL is also authorized in Moldova to:

- register, notify, renew or modify the registration of the products as listed in Appendix A;
- participate in public tenders for supply of automated clinical chemistry analyzers;
- perform service activities.

We guarantee that the quality of our products is corresponding to the requirements for IVD products.

Products will be invoiced via: **ELITech Clinical Systems SAS** Zone Industrielle 61500 Sées France

This Manufacturer Authorization Letter (MAL) is governed by and construed in accordance with Dutch law and is valid for a period of two (2) years unless terminated with a written notice by the issuer.

ELITechGroup B.V.

Maurice Verdaasdonk Managing Director ELITechGroup B.V. P.O. Box 100 – 6950 AC Dieren Van Rensselaerweg 4 – 6956 AV Spankereiv The Netherlands





Appendix A - List of products

• SELECTRA MACH5 Including all accessories and parts





ELITechGroup B.V. P.O.Box 100 6950 AC Dieren Van Rensselaerweg 4 6956 AV Spankeren The Netherlands T: +31 313 430 500 F: +31 313 427 807 info.ecsnl@elitechgroup.com www.elitechgroup.com Chamber of Commerce 09175642

To: Whom it May Concern

Regulatory status of parts & accessories

As mentioned on the current Declarations of Conformity of our Clinical Chemistry Analyzers also the accessories conform to the provisions of the EU Directive on In Vitro Diagnostic Medical Devices (98/79/EC). This applies to the parts and accessories as mentioned in the attached list.

'IVD accessory' means an article which, whilst not being an IVD medical device, is intended specifically by its manufacturer to be used together with an IVD device to enable that IVD device to be used in accordance with its intended purpose.

ELITechGroup B.V.

Adriaan P. Intveld Manager Quality Assurance & Regulatory Affairs





Part number	Description	IVD medical device	IVD accessory	general laboratory use	spare part	supporting part
1540-001	Anti-Slip sheet					✓
2206-007	Cooling Liquid (1 L)					✓
3062-021	Sample cup (1000 pcs)		\checkmark			
3062-033	Sample tube 6 ml (500 pcs)					✓
3062-040	Water container 10 L					✓
3062-041	Water container 5 L					✓
3066-155	Syringe 100 μl		\checkmark			
3066-156	Syringe 1 ml		\checkmark			
3069-040	Keyboard Dust cover					✓
3069-047	Keyboard Dust cover					✓
3070-518	Cap holder					✓
3070-538	Cap rotor Left					✓
3070-539	Cap rotor right					✓
3201-002	Dichromate 8 Abs (25ml)		\checkmark			
3365-192	USB Stick					✓
3374-003	Mains cable (USA)					✓
3374-059	Pumpunit cable		✓			
3374-066	Mains cable					✓
3374-097	Serial Null-modem cable					✓
3374-286	USB Extension cable					✓
4804-038	Reagent identification Disc					✓
6001-826	Diluted Waste container		✓			
6001-827	Concentrated Waste container		✓			
6001-860	Water container		✓			
6001-861	Tube assy (analyser)		✓			
6001-872	Tube assy (cooling unit)		✓			
6002-102	Assorter unit				✓	
6002-386	System software on CD		✓			
6002-706	Reaction Rotor set (3 pcs)		✓			
6002-726	System Disc		✓			
6002-817	Bottle 30 ml (20 pcs)		✓			
6002-818	Bottle 15 ml (20 pcs)		✓			
6002-904	Water container 5 L		✓			
6002-910	Assorter unit				✓	
6002-913	External tubing		✓			
6003-074	System software on USB stick		✓			
6003-444	Diluted Waste Container 5 L		✓			
6003-466	Keyboard Support option					✓
6003-797	CW Waste Container 2 L		\checkmark			1
6003-808	Assorter unit				\checkmark	1







Zone Industrielle – 61500 SEES – France Tél. : + 33 (0)2 33 81 21 00 / Fax : + 33 (0)2 33 28 77 51

REAGENTS

TO WHOM TO BE CONCERNED

We, Seppim S.A.S., manufacturers of Elitech Clinical Systems reagents, having our factory at Zone Industrielle, 61500 Sées - France, confirm that our clinical reagents have been validated on Vital Scientific equipment. As such available Elitech Clinical Systems reagent applications for Vital Scientific instruments are CE-IVD compliant.

Reagents, other than Elitech Clinical Systems reagents, are not validated on Vital Scientific equipments, and we also can't know the impact of other reagents on Vital Scientific equipments.

May 22nd, 2012

Noi, subsemnații Seppim S.A.S., compania producătoare a regenților Elitech Clinical Systems, având fabrica de producere în Zone Industrielle, 61500, Franța, confirmăm, că reagenții au fost testați si validați pe echipamentele Vital Scientific. Pentru acești reagenți existând și protocoale specializate pentru analizatoarele produse de Vital Scientific. Atât reagenții cât și echipamentele sunt certificate CE-IVD.

Alți reagenți înafara de Elitech Clinical Systems, nu au fost testați și validați la echipamentele Vital Scientific și noi nu cunoaștem compatibilitatea și impactul lor asupra analizatoarelor Vital Scientific.

22 mai 2012

Signed on behalf of the manufacturer Valérie GOURDON Regulatory Affairs Manager COMPANY SEPPIM S.A.S

> SEPPIM S.A.S 4 rue Auguste Mottin Zone Industrielle 61500 SEES – FRANCE Tél. +33 (0)2 33 81 21 00 - Fax +33 (0)2 33 28 77 51 SIREF : 318 365 228 00036

Société par actions simplifiée au Capital de 1 219 592.14 € SIRET 318 365 228 00036 APE 2059Z RC ALENCON 318 365 228