

00-04/21

REPROCESSING INSTRUCTIONS

## CE

## Reprocessing of resterilizable medical devices with assembly instructions in accordance with DIN EN ISO 17664:2018 **Risk assessment groups Critical B**

Manufacturer:	FEHLING INSTRUMENTS GmbH & Co. KG			
Products:	All instruments or medical devices of the above mentioned risk assessment grou supplied by FEHLING INSTRUMENTS GmbH & Co. KG, for which the following additional assembly instructions are available:			
	Flexible holding device MNU-1V			
	Bar retractor with pinion			
	Bar retractor with sprocket/lockM30			
	Vascular retractor			
	LEYLA titanium holding device			
	Operating table adapting clamp			
	Handle for CERAMO <sup>®</sup> FI Micro Set			
	Blade guides (for ball adapters) and guide forceps			
Warnings:	Do not clean CERAMO <sup>®</sup> instruments (recognizable by their black-brown surfa and titanium instruments with oxidative processes (processes using hydro peroxide H <sub>2</sub> O <sub>2</sub> , e.g. Orthovario or Oxivario from Miele). By dissolving titanium application of these procedures leads to the destruction of titanium instrument the titanium-containing CERAMO <sup>®</sup> coating after some time.			
	Similarly, instruments with components of plastics should not be cleaned			
	oxidative processes. These processes lead to thermal-oxidative aging of material, which may under certain circumstances not be detectable by vis discoloration or embrittlement.			
	The instruments may only be used, reprocessed and disposed of by qual medical personnel.			
	Handle instruments with care during storage, transport and cleaning! Avoid stri and applying pressure to instruments, so as not to cause any conseque damage! Do not overstrain functional parts!			
	The medical device is to be reprocessed prior to use. The instrument must under risk assessment according to the RKI Guidelines (non-critical, semi-critical, critical, A/B/C) prior to reprocessing.			
	The national legal regulations, national and international standards and guidel as well as the company's own hygiene regulations for reprocessing are to complied with.			
	The applicable national regulations must be followed for the reprocessing instruments used in patients with Creutzfeldt-Jakob disease (CJD), susper CJD or possible variants.			
	SUPERPLAST instruments:			
	Thermal disinfection and steam sterilization should be used to activate the sh memory. The following should be observed here:			
	<ul> <li>SUPERPLAST instruments must be stored in such a way that they are prevented from regaining their original shape by environmental influer (e.g., other instruments or restricted space).</li> </ul>			
	<ul> <li>After disinfection/sterilization, allow the SUPERPLAST instruments to down to room temperature. The functionality of the instruments may impaired if they are bent at temperatures in excess of 40°C.</li> </ul>			
Limitations on reprocessing:	Frequent reprocessing has little impact on these instruments. The end of produlifie is normally determined by wear and tear and damage occurring through us (e.g. damage, illegible marking, functional failure - also see "Maintenanc Checking and Testing").			



R 16 EN

00-04/21

REPROCESSING INSTRUCTIONS

## CE

Instructions			
General information on reprocessing:	Reprocessing is based on a validated procedure. All the cleaning steps mentioned (manual pre-cleaning, automated/manual cleaning, manual disinfection, and sterilization) were validated with the respective parameters specified in each case and listed under "Validated procedure". The recommended reprocessing agents (detergent: Neodisher® MediClean forte (Dr. Weigert); disinfectant: Korsolex® med AF (Bode Chemie GmbH)) were used for validation. Both water of potable water quality and fully deionized water (deionized water, demineralized, microbiologically at least of potable water quality) are used for cleaning. Automated reprocessing is preferable to manual cleaning due to a better and safer cleaning result. There is also the option of cleaning our instruments with other tested and approved chemicals which have been recommended by the chemical manufacturer with regard to their material compatibility. Please always observe the manufacturer's information on concentration, exposure time, temperature and renewal of the detergents and disinfectants. All instructions for use of the chemical manufacturer must be strictly observed. Otherwise, this can lead to visual material changes or material damage, such as, for example, corrosion, fractures or premature aging.		
Initial treatment at the place of use:	Pre-cleaning: ensure that blood, tissue and drug residues are removed from the instruments with a disposable cloth/paper wipe immediately after completion of the procedure and that they undergo mechanical cleaning immediately. After completion of initial treatment of the instruments, visual inspections must be performed to ensure that the instruments are complete. The instruments must be transported from the place of use to the place of reprocessing such that neither the user, third parties, the environment nor the medical devices are endangered or damaged (placement in closed, puncture-proof containers and - if necessary - use of protective caps).		
Preparation prior to cleaning:	It is recommended to reprocess the instruments immediately after use because it is very difficult to remove dried residues from instrument parts that are difficult to access. Do not immerse in normal saline solutions (risk of pitting or stress corrosion cracking). Instruments which were connected to each other during use must be disassembled again before cleaning. Then disassemble take-apart instruments according to the corresponding assembly instructions.		
Manual pre-cleaning	Validated procedure:         Equipment:       Basin         Soft brush       Water spray gun (or similar)         Detergent:       Neodisher® MediClean forte (Dr. Weigert)         Procedure/Parameters:       •         •       Rinse instruments, if possible in disassembled condition, under running cold water of potable water quality (<40°C) until all visible contamination has been removed. Remove stubborn contamination with a soft brush (not a wire brush!).		

## FEHLING R 16 EN INSTRUMENTS 00-04/21

REPROCESSING INSTRUCTIONS

CE

	<ul> <li>If necessary, the moving parts of the instrument are moved back and for the cleaning bath.</li> </ul>		
	• During the exposure time, use appropriate brush (not a wire brush) to remove coarse contamination.		
	• Rinse the instruments for one minute in cold deionized water (see "Gener Information on Reprocessing") and, if applicable, move movable parts bac and forth.		
Cleaning/Disinfection:	If possible, a washer/disinfector according to DIN EN ISO 15883, which uses thermal disinfection, is to be preferred.		
Cleaning: Automated	Avoid overfilling instrument trays and washing trays - use only suitable instrume holders. When placing instruments in the sterilization baskets and removing the afterwards, take special precautions to ensure that the tips do not become stud in the mesh.		
	Validated precedure:		
	Validated procedure: Equipment: Washer-disinfector		
	G 7835 CD (Miele) / PG 8535 (Miele)		
	Cleaning program: Des-Var-TD (G 7835 CD)		
	Detergent: Neodisher <sup>®</sup> MediClean forte (Dr. Weigert)		
	<ul> <li>Preparation:</li> <li>Instruments with joints are to be placed in the device such, that the joints a opened or disassembled if possible, and that the water can flow from the cavities and sac holes.</li> <li>If applicable, loosen springs</li> <li>Ensure that the inside of all cavities is also completely rinsed.</li> <li>Ensure that no areas are left unwashed.</li> <li>Connect the Luer connections of the instruments, if present, to the Luer Lo flushing attachment of the WD.</li> </ul>		
	Procedure/Parameters:		
	<ul> <li>Pre-wash for 3 minutes with cold water (potable water quality, &lt;40°C)</li> <li>Emptying</li> </ul>		
	<ul> <li>Clean for 10 minutes with a solution of 0.5 - 2% Neodisher<sup>®</sup> MediClean for in water (potable water quality) at 55°C</li> <li>Emptying</li> </ul>		
	<ul> <li>Rinse for 2 minutes with water (potable water quality, &lt;40°C)</li> </ul>		
	Emptying		
	Rinse for 1 minute with cold deionized water (<30°C)		
	• Emptying		
	<ul> <li>Thermodisinfection for 5 minutes with deionized water (&gt;90°C)</li> <li>Dry for 30 minutes (90°C)</li> </ul>		
	After cleaning in the machine, inspect cavities, blind holes, etc. for visib contamination. If necessary, repeat the cycle or clean manually.		



R 16 EN

00-04/21

REPROCESSING INSTRUCTIONS CE

Cleaning: Manually	Validated procedure:		
0 ,	Equipment:	Basin	
		Soft brush	
		Water spray gun (or similar)	
		Bandelin Sonorex Digitec	
	Detergent:	Neodisher <sup>®</sup> MediClean forte (Dr. Weigert)	
	Procedure/Parameter	<u>S:</u>	
		s, if possible in disassembled condition, in cold water (potable °C) for 10 minutes.	
	• Move any movable parts, if present, back and forth over the entire range of movement.		
	<ul> <li>Use a soft brush (not a wire brush) to clean the instruments unt contamination is visible.</li> </ul>		
	• Rinse the instruments for at least 20 seconds with a water spra similar).		
	<u>Ultrasonic cleaning:</u>		
	<ul> <li>Clean for 10 minutes at &lt; 40 °C with 0.5 - 2% cleaning solution at 3</li> </ul>		
	• After ultrasonic cleaning, rinse the instruments with a water spray gun (or similar) for at least 20 seconds.		
	• Rinse the instrum <40°C).	ents for at least 10 seconds with water (potable water quality,	
		<40°C) is to be used for the final rinse. Rinse the instruments ater for at least 30 seconds. Ensure that no residues remain	
Disinfection: Manually	Consult the instructions on the label when selecting a disinfectant (se manufacturer information).		
	Validated procedure:		
	Validated procedure: Equipment:	Basin	
	Equipment.	Bandelin Sonorex Digitec	
	Disinfectant:	Korsolex <sup>®</sup> med AF (Bode Chemie GmbH)	
	Procedure/Parameter	s:	
	After cleaning, pla suitable disinfecta Ensure that all su	ace the products in an ultrasonic bath (35 kHz, <40°C) with a ant solution (e.g. 0.5% Korsolex <sup>®</sup> med AF) for 5 minutes. Infaces are wetted with the disinfectant. If applicable, move in the disinfection bath before switching on the ultrasonic	
	for at least 1 min	rinse all products thoroughly with deionized water (<40°C) ute to remove the disinfectant and, if applicable, move the f the instrument back and forth.	
	Ensure that no re	sidues remain on the products.	
	• Dry with sterile, o	il-free compressed air.	
Drying:	120°C. Then dry with	eved as part of the cleaning/disinfection cycle, do not exceed a suitable compressed air in accordance with Robert Koch nendations. Pay particular attention to the drying of difficult-	

FEHLING INSTRUMENTS

00-04/21

R 16 EN

REPROCESSING INSTRUCTIONS

Maintenance, checking	Assemble the disassembled instruments according to the corresponding as		
and testing:	<ul> <li>instructions.</li> <li>For instruments with movable components that are exposed to friction (e.g. joints), an instrument oil based on paraffin/white oil (according to the valid European or United States Pharmacopoeias) which is biocompatible, steam sterilizable and steam-permeable is to be applied. Such places are additionally marked by a corresponding symbol of an oil can. Instruments must not be treated with care products containing silicone. These can lead to stiffness and question the effect of steam sterilization.</li> </ul>		
		e instruments prior to each use. When doing so, check fractures and mechanical malfunctions and missing	
	Check instruments with movable parts for smooth operation (avoid excest play). Check locking mechanisms. All instruments: use a magnifying lamp to visually inspect the components damage and wear and tear.		
	In particular, inspect the crit Defective or damaged instru- out and cleaned and disir Repairs may only be carried by the manufacturer. A ve manufacturer.	ical points on moving parts and in the working area. Iments, or those with illegible markings, must be sorted infected before being returned to the manufacturer. I out by the manufacturer or by workshops authorized rification form for this process is available from the	
	accordance with hospital pr sharp edges in particular,	ger be repaired must be disposed of as scrap metal in actice. In the case of surgical instruments with tips or safe storage in a closed, puncture and break-proof be ensured. Do not use damaged instruments!	
Packaging:	Singly: In accordance with the standard series DIN EN 868, DIN EN ISO 11607, and DIN 58953. Sets: sort instruments into dedicated trays or place them in general-purpose sterilization trays. Pack the trays appropriately using a suitable procedure.		
Sterilization:	Steam sterilization in a fractionated vacuum process in a device complying with DIN EN 285 and DIN EN ISO 17665. In order to prevent staining and corrosion, the steam must be free of contaminants. The recommended limits for contaminants for feed water and steam condensate are defined by DIN EN 285.		
	Validated procedure:		
	Equipment:	Tuttnauer Autoclave Type B 3870 EHS / Lautenschläger ZentraCert	
	Procedure/Parameters: Cycle type: Sterilization temperature: Holding time:	3 pre-vacuum phases 132 – 134°C 4 – 5 minutes	
	Drying time:	20 minutes	
		one instrument in a sterilization cycle, do not exceed erilizer (see manufacturer's instructions).	



R 16 EN

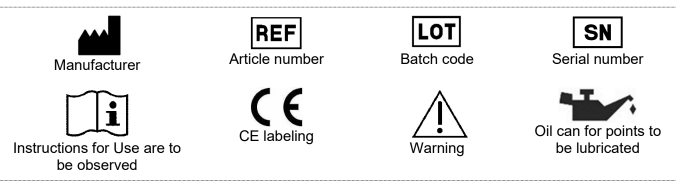
00-04/21

REPROCESSING INSTRUCTIONS CE

Storage:	In accordance with §4 of the German Medical Devices Operator Ordinance (MPBetreibV) and standard series DIN EN 868, DIN EN ISO 11607, and DIN 58953. Instruments must be stored dry, at room temperature, clean, protected from damage and mechanical influences (avoid condensation, damage). Always keep instruments, if applicable, in a released state. This counteracts premature fatigue of the spring tension. Instruments must be transported to their place of use in a closed, puncture-proof sterile container.
Disposal:	These products largely consist of steel or titanium. These are to be cleaned prior to disposal. Disposal can be performed at a scrap metal recycling facility. To protect employees, care must be taken to ensure that any pointed tips or sharp edges are protected.
Obligation to report serious incidents:	The user is obliged to report serious incidents relating to the medical device to the manufacturer per e-mail to vigilance@fehling-instruments.de or via the report form at https://www.fehling-instruments.de/en/complaint/ and the competent authority of the Member State where the user is registered.
To contact the manufacturer:	FEHLING INSTRUMENTS GmbH & Co. KG Hanauer Landstr. 7A 63791 Karlstein, Germany Tel.: +49 (0) 6188-9574-40 Fax: +49 (0) 6188-9574-45 E-mail: info@fehling-instruments.de www.fehling-instruments.de

**Symbols** 

In as far as the medical device or medical device label or reprocessing instructions are labeled, the symbols represent the following meaning:



Any modification to the device or any deviation from these Instructions for Use will result in exclusion of liability! Subject to change without notice.

The instructions listed above were validated by the medical device manufacturer as suitable for preparing a medical device for reuse. It is the responsibility of the reprocessor to ensure that the reprocessing actually performed using equipment, materials, and personnel in the reprocessing facility achieves the desired results. This normally requires validation and routine monitoring of the process. Likewise, any deviation from the provided instructions on the part of the reprocessor should be properly evaluated for effectiveness and potential adverse consequences.