# | HEINE BETA® 200 LED retinoscope.



DATA		
Description	HEINE BETA 200 LED retinoscope 2,5 V	HEINE BETA 200 LED retinoscope 3,5 V
Catalogue number	see catalogue or price list	

Document release date June, 2024

### MECHANICAL

Weight	90 g
Weight packaging (including product)	200 g
Dimensions product	129 x 37 x 43 mm
Dimensions packaging	Case: 158 x 90 x 43 mm  Hard case BETA 200 LED retinoscope: 186 x 120 x 50 mm  Hard case BETA 200 LED diagnostic set: 237 x 138 x 52 mm
Connections	AV connection
Imprints	Product name, ParaStop, HEINE made in Germany, CE, HEINE logo, datamatrix code, serial number, www.heine.com

#### GENERAL

Materials	metal, glass, plastic			
REACH   RoHS	conform			
Biocompatibility	conform			
Surface	metal, glass, plastic			
Envrionmental conditions operation	Temp.: +10°C to +35°C, relative humidity: 30	Temp.: +10°C to +35°C, relative humidity: 30%-90%, air pressure: 800hPa to 1060hPa		
Envrionmental conditions storage	Temp.: -10°C to +55°C, relative humidity: 10	Temp.: -10°C to +55°C, relative humidity: 10%-95%, air pressure: 700hPa to 1060hPa		
Envrionmental conditions transport	Temp.: -40°C to +70°C, relative humidity: 10%-95%, air pressure: 500hPa to 1060hPa			
Durability	5 years warranty			
Instructions for use*	Deutsch, English, Français, Español, Italiano, Svenska, Nederlands, Dansk, Norsk, Suomi, Português			
Operating elements	Light intensity control, single control for vergence and rotation, ParaStop for precise and easy selection of a parallel beam, detachable brow rest			
Power supply	HEINE battery handles (2,5 V)  HEINE rechargeable handles (3,5 V), HEINE EN 200 wall transformator			
Accessories	Fixation cards			

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Input voltage	1,8 V-3,2 V	3,0 V-3,7 V
Power consumption	typ. 373 mA at full brightness and 3,2 V	max. 400 mA typ. at full brightness and nominal supply
Protection class	internally powered	charging: II, operating: internally powered

## OPTICAL

Туре	LED (HQ) illumination (2,5 V   3,5 V)
Light controlling	Rheostat (continuous brightness control)
Color temperature	typ. 3000 K
Length of the streak (500 mm distance)	typ. 35 mm
Width of the streak (500 mm distance)	typ. 1,1 mm
Working distance	500 mm
Medium life expectancy (LED)	> 50.000 h
Classification according to ISO 15004-2	Group 2

# HYGIENIC REPROCESSING

Procedure	Please see detailed description for the reprocessing procedure online at www.heine.com
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## CODES

Customs code (tariff number)	90185090
GTIN	4053755119967 (2,5 V); 4053755191598 (3,5 V)
Traceability	UDI-Code
Country of origin	Germany (DE)

## REGULATORY

Product classification (EU)	Class I
Product classification (USA)	Class 1, 510(k) exempt
Product classification (Canada)	Class I
UMDNS code	13-372
GMDN code	32712
Regulation number (FDA)	886.1780
Product code (FDA)	HKM

#### FULFILLS THE REQUIREMENTS OF DIRECTIVES & STANDARDS

ISO 13485	Medical devices - quality management systems - requirements for regulatory purposes		
Regulation (EU) 2017/745	European regulation for medical devices (MDR)		
IEC 60601-1	Medical electrical equipment: general requirements for basic safety and essential performance		
IEC 60601-1-2	Medical electrical equipment - part 1-2: general requirements for basic safety and essential performance - collateral standard: electromagnetic disturbances - requirements and tests		
ISO 14971	Medical devices - application of risk management to medical devices		
IEC 60601-1-6	Medical electrical equipment - part 1-6: general requirements for basic safety and essential performance - collateral standard: usability		
IEC 62366-1	Medical devices - part 1: application of usability engineering to medical devices		
ISO 15004-1	Ophthalmic instruments - fundamental requirements and test methods - part 1: general requirements applicable to all ophthalmic instruments		
ISO 15004-2	Ophthalmic instruments - fundamental requirements and test methods - part 2: light hazard protection		
ANSI Z80.36	Ophthalmics - light hazard protection for ophthalmic instruments		
ISO 12865	Ophthalmic instruments - retinoscopes		
IEC 60601-1-9	Medical electrical equipment - part 1-9: general requirements for basic safety and essential performance - collateral standard: requirements for environmentally conscious design		
ISO 10993-1	Biological evaluation of medical devices - part 1: evaluation and testing within a risk management process		
ISO 17664	Processing of health care products - information to be provided by the medical device manufacturer for the processing of medical devices		
ISO 2248	Packaging; complete, filled transport packages, vertical impact test by dropping		
Directive (2011/65/EU) ROHS	On the restriction of the use of certain hazardous substances in electrical and electronic equipment		
Directive (2012/19/EU) WEEE	On the waste electrical and electronic equipment		
Regulation (1907/2006) REACH	Registration, evaluation, authorization and restriction of chemicals		
Directive (94/62/EC) packaging packaging waste	Packaging and packaging waste, German registration no. DE 5329703000126		

<sup>\*)</sup> further languages on request

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