

AN ITALIAN COMPANY THAT RESEARCHES, DEVELOPS AND PRODUCES READY TO USE KITS USING CHROMATOGRAPHIC TECHNIQUES FOR CLINICAL LABORATORIES

CONTROL LYOPHYL. IN URINE FOR 3-METHYLINDOLE - LEVELS 1 and 2 Code Z60019

OBJECTIVE

These controls in plasma are used for the internal quality control and serve to monitor the accuracy and precision of analytical procedures devoted to the quantitative determination of analytes contained therein. These controls are lyophilized human matrix and are available in two different concentration ranges. Should be handled as if they were a real patient sample.

RECOVERY

Remove the metal seal and rubber stopper from the vial. Add exactly 1 ml of HPLC-grade water through the tube. Replace the rubber stopper, allow to stand for at least 5 to 10 minutes. Before use, mix by inverting the vial to dissolve the material until a homogeneous solution.

STORAGE AND STABILITY

The controls are stable for 12 months from the date of preparation if stored at -20 ° C. After reconstitution are stable for 1 day at -20 °C. Do not use after the expiry date.

PRECAUTIONS

These controls in human matrix should be treated with care and treated as potentially infectious.

LO1		M	\square	
063		Oct	Oct	
		2023	2024	

PACKAGINGS AVAILABLE:

- Z60019 CONTROL IN URINE FOR 3-METHYLINDOLE (Skatole) - LEVELS 1 and 2 2 x 5 x 1 ml

Release N° 002	September 2019



ANALYTE	MEASUREMENT UNIT	MEDIUM VALUE L1	RANGE L1	MEDIUM VALUE L2	RANGE L2
3-METHYLINDOLE	μg/l	15,3	10,7 – 19,9	75,4	52,8 - 98,1

This product fulfills all the requirements of Directive 98/79/EC of 27/10/1998 on in vitro diagnostic medical devices (IVD). The declaration of conformity is available upon request.

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FOR IN VITRO DIAGNOSTIC USE ONLY

produced by

COMPANY WITH
MANAGEMENT SYSTEM
CERTIFIED BY DNV
= ISO 9001=
= ISO 13485 =



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