

RANDOX

CERTIFICATE OF ANALYSIS

This is to certify that the following kit was tested by a
Randox Testing Laboratory and approved by the QC Department.

DATE OF MANUFACTURE: 2025-08-29
KIT DESCRIPTION: Immunoassay Premium Plus Level 3(IA
PREMIUM PLUS 3)
CATALOGUE NO: IA3111
LOT NUMBERS: 2653EC
EXPIRY DATE: 2028-08-28
SYSTEM OF ANALYSIS: Roche e801

TEST CONTROLS OR VIALS

ANALYTE	LOT NUMBER	RANGE	TARGET	RESULT	PRECISION (if applicable)
FSH mU/mL	2653EC	44.4-66.6	55.5	55.0	N/A
	2653EC	44.4-66.6	55.5	54.5	N/A
LH mU/mL	2653EC	41.8-62.8	52.3	55.2	N/A
	2653EC	41.8-62.8	52.3	54.5	N/A
Progesterone ng/mL	2653EC	22.8-34.2	28.5	28.7	N/A
	2653EC	22.8-34.2	28.5	27.6	N/A

QC TESTED BY: MAYA PULIKUNNI VIJAYAN
DATE: 06-Nov-25
RESULT (PASS/FAIL): Pass
QC APPROVED BY: *A. Martin*

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