

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: November 2016
KIT DESCRIPTION: IA PREMIUM TRI - LEVEL
CATALOGUE NO: IA2633
LOT NUMBERS: 402366 (1625EC, 1626EC, 1627EC)
EXPIRY DATE: 2019-04-28
SYSTEM OF ANALYSIS: Siemens Immulite 2000 Xpi

TEST CONTROLS OR VIALS

ANALYTE	LOT NUMBER	RANGE	TARGET	RESULT	PRECISION (if applicable)
CEA (ug/l)	1625EC	3.82 - 5.72	4.77	3.8	
				4.0	
	1626EC	19.1 - 28.7	23.9	21.6	
				20.7	
	1627EC	59.8 - 89.6	74.7	70.9	
				71.5	
DHEAS (umol/l)	1625EC	1.64 - 2.46	2.05	1.8	
				1.8	
	1626EC	9.6 - 14.4	12.0	11.7	
				11.3	
	1627EC	16.4 - 24.6	20.5	20.4	
				19.5	
Alpha Foetal Protein (kU/L)	1625EC	8.16 - 12.2	10.2	9.2	
				9.3	
	1626EC	42.9 - 64.3	53.6	53.2	
				51.9	
	1627EC	157 - 235	196	202.0	
				184.0	

QC TESTED BY: Davina Graham
DATE: 11 Nov 16
RESULT (PASS/FAIL): pass
QC APPROVED BY: *Adelle Martin*

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