Ultrasensitive PSA (UPSA)

Accreditation Status	UKAS Schedule of Accreditation	
Date Scheme started:	2019	
Clinical Applicability:	A marker of recurrence for post radical prostatectomy patients	
Analytes:	UPSA (SER/058)	
Units for Reporting:	$\mu\text{g/L}$ in relation to the WHO International Standard	
Samples Distributed:	Liquid format. Normal and pathological human serum	
Number of Distributions per year:	12	
Number of Samples per Distribution:	2	
Frequency of Distributions:	Every month as outlined in the Distribution Schedule	
Schedule of Analysis:	Data entry is via the web for the submission of results. Data analysis is commenced 14 days after sample dispatch. Late returns are accepted and will contribute to the laboratory's cumulative performance statistics	
Data Analysis:	All Laboratory Trimmed Mean (ALTM) with truncation at 2SD, SD, and CV%. Reports also show method specific statistics. Individual laboratory performance is expressed in terms of MRBIS, SDBIS, and MRVIS	
	Chosen Coefficient of Variation for ultrasensitive Prostate Specific Antigen is 12.5%	
Performance Scoring:	MRVIS	
Criteria of Performance:	Laboratory performance for ultrasensitive PSA is classified in terms of the MRVIS over a running analytical window of 12 Distributions (12 months)	
Persistent Poor Performance:	Ideal MRVIS Good Adequate Poor Defined as being in the Poor Performance category	<50 50 - 100 101 - 200 >200 or SDBIS >200
	successive Distributions	