

## Pilot Point of Care C-Reactive Protein (CRP) Testing

<b>Accreditation Status:</b>	<i>currently not accredited to ISO 17043:2010</i>		
<b>Date Scheme started:</b>	2017		
<b>Clinical Applicability:</b>	Monitoring of the acute phase response		
<b>Analytes:</b>	C-Reactive Protein ( <b>POC/555</b> )		
<b>Units for Reporting:</b>	mg/L		
<b>Samples Distributed:</b>	Liquid format. Normal and pathological human serum		
	Additional materials may be produced for specific recovery experiments by the addition of purified CRP to an analyte-free serum matrix		
<b>Number of Distributions per year:</b>	4		
<b>Number of Samples per Distribution:</b>	2		
<b>Frequency of Distributions:</b>	Currently seasonal as outlined in the <a href="#">Distribution Schedule</a>		
<b>Schedule of Analysis:</b>	<a href="#">Data entry</a> 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		
<b>Data Analysis:</b>	All Laboratory Trimmed Mean (ALTM) with truncation at 2SD, 3SD, 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 C-Reactive protein is 8%		
<b>Performance Scoring:</b>	MRVIS		
<b>Criteria of Performance:</b>	Laboratory performance is classified in terms of the MRVIS over a running analytical window of 4 Distributions (12 months)		
	Ideal	MRVIS	<50
	Good		50 - 100
	Adequate		101 - 200
	Poor		>200 or SDBIS >200
<b>Persistent Poor Performance:</b>	Defined as being in the Poor Performance category for two or more successive Distributions		