



SCOPE OF ACCREDITATION TO ISO/IEC 17043:2010

CANADIAN MICROBIOLOGY PROFICIENCY TESTING (CMPT)
Department of Pathology and Laboratory Medicine/University of British Columbia
G408 - 2211 Wesbrook Mall, Vancouver, British Columbia V6T 2B5, Canada
Dr. Lucy Perrone, Chair, CMPT
Phone: 604 827 1337
Email: cmpt.path@ubc.ca

PROFICIENCY TESTING PROVIDER

Valid To: May 31, 2027

Certificate Number: 3749.01

In recognition of the successful completion of the A2LA evaluation process, this proficiency testing provider has been found to meet ISO/IEC 17043:2010, "Conformity assessment – general requirements for proficiency testing".

Accreditation is granted to this provider to provide proficiency testing schemes in the following areas:

<u>PROGRAM NAME*</u>	<u>SCHEME DESCRIPTION</u>
Clinical Bacteriology ^{1,2,3} 4 shipments /year	<p>Laboratories participating in the clinical bacteriology CMPT program are separated into peer categories (A and C1) based on a recommended program-developed work-related weighted rating scale. The rating scale is based on complexity of samples processed. Individual laboratories and provincial accreditation programs decide 'best fit' for the participating laboratories. Shipments consist of up to 7 challenges (5 simulated samples, including a Gram smear ± pre- or post-analytical scenario) depending on the category of the laboratory.</p> <p><u>Category A Laboratories:</u> Large tertiary-care laboratories that perform most clinical testing, including blood cultures, fluids, CSF, tissues, and other critical specimens. Only Category A laboratories receive all available types of samples.</p> <p><u>Category C1 Laboratories:</u> Set-up facilities only. May perform Gram staining and address pre-analytic issues.</p> <p>The types of samples and bacterial isolates to be shipped are selected by the CMPT committee annually and each shipment consists of simulated clinical samples, which laboratories may receive routinely. Participants are required to report the cellular and bacterial components of the Gram smear, correctly identify the pathogens, if present, and/or report the related susceptibilities. For the pre- or post- analytical challenges, participants are given a laboratory scenario and they must choose the best answer from a list of responses provided.</p>
<i>Clostridioides difficile</i> assay ^{1,2} 2 shipments /year	<p>The <i>Clostridioides difficile</i> program is designed for the detection of common antigen and/or toxin in stool samples. The samples can be tested by culture, enzyme immunoassay and molecular methods.</p> <p>The program consists of 2 simulated stool samples. Participants are required to report "positive" or "negative" for the presence of the common antigen and/or toxin.</p>

<u>PROGRAM NAME*</u>	<u>SCHEME DESCRIPTION</u>
Supplementary Gram Smears^{1,2} 2 shipments /year	The Supplementary Gram Smear program consists of 2 slides for Gram staining (2 simulated fluid samples, usually cerebrospinal, joint fluid gram smear). Participants are required to report the cellular and bacterial components of the slides.
Enteric Parasitology^{1,2} 3 shipments /year	This program is intended for laboratories performing enteric parasitology tests by conventional microscopic analysis. Shipments consists of 3 SAF (sodium acetate-acetic acid-formalin) preserved stool samples. Participants are required to report any helminths and/or protozoa present in the concentrate and/or the direct, stained smear.
Mycology Plus¹ 3 shipments /year	This program consists of 3 Mycology samples (consisting of yeasts, dermatophytes, molds, fungi) and 3 fungal smears. Participants are required to report the identity of the yeast, dermatophyte, mold and/or fungus in each of the samples and/or report any applicable anti-fungal susceptibilities. Participants are also required to report “positive” or “negative” for the presence of hyphae/pseudohyphae in the fungal smears.
Water Microbiology³ 3 shipments /year	The CMPT Water Microbiology Program is intended primarily for laboratories performing drinking/recreational water assessments. Samples are simulated stabilized waters that can be assessed by using membrane filtration, enzyme substrate, MPN, and Presence/Absence methods. <u>Drinking Water Program:</u> Participants are required to identify and enumerate total coliforms, thermotolerant coliforms and <i>E.coli</i> and Heterotrophic Plate Counts, in drinking water samples. Shipments consist of 4 samples. Testing methods used by the participants include Membrane Filtration, Enzyme Substrate, Most Probable Number (MPN), Presence/Absence and Heterotrophic Plate Count (HPC). <u>Recreational Water Program:</u> Participants are required to identify and enumerate <i>Enterococcus faecalis</i> in simulated marine water samples, <i>Escherichia coli</i> in freshwater beach samples and <i>Pseudomonas aeruginosa</i> in spa/swimming pool water samples. Shipments consist of 3 samples. Testing methods used by the participants are membrane filtration and enzyme substrate. <u>Heterotrophic Plate Count:</u> Participants are required to enumerate <i>Escherichia coli</i> and <i>Enterobacter</i> species in simulated drinking water samples. Shipments consist of 3 samples. Testing methods used by the participants are membrane filtration and enzyme substrate.
Soil/Sludge^{1,3} 2 shipments /year	This program is intended primarily for environmental laboratories performing soil/sludge testing for <i>Salmonella</i> species. Samples are simulated soil sludge matrices that can be assessed by using membrane filtration, enzyme substrate, MPN, and Presence/Absence methods. Labs must quantify the CFU's. Each shipment consists of 2 samples.

<u>PROGRAM NAME*</u>	<u>SCHEME DESCRIPTION</u>
Trichomonas vaginalis Antigen¹ 3 shipments /year	The program's objective is to provide EQA to those laboratories performing laboratory diagnosis of trichomoniasis using the OSOM <i>Trichomonas</i> Rapid Test (Sekisui Chemical Co) or molecular methods. Each shipment has 4 samples. Participants are required to report if the samples are "positive" or "negative" for <i>Trichomonas vaginalis</i> antigen.
Screening and Molecular¹ 2 shipments /year	The program is designed for laboratories that use screening (eg. Chromogenic agar) molecular methods to detect the presence of MRSA, VRE, CRE and group B <i>streptococcus</i> . Four samples of each type are sent in each shipment. Participants are required report "positive" or "negative" for each of the 3 types of samples.
Shiga Toxin¹ 2 shipments /year	The program's objective is to provide EQA to those laboratories performing testing to detect organisms producing Shiga toxin. Three simulated stools samples are in each shipment. Methods that can be used are culture, toxin or molecular methods. Participants are required to report "positive" or "negative" for the presence of Shiga toxin.
Acid-Fast Bacilli¹ 2 shipments /year	The program's objective is to provide EQA to those laboratories performing testing to detect organisms that are acid-fast. Acid-fast stains are used so that bacilli, if present, that can resist the acid in the method, can be observed microscopically. Each shipment has 3 samples. Participants are required to report "positive" or "negative" for the presence of acid-fast bacilli.
Enteric Panel¹ 2 shipments /year	The program's objective is to provide EQA to those laboratories performing testing to detect the presence of <i>Salmonella</i> , <i>Shigella</i> species, toxigenic <i>E.coli</i> , <i>Yersinia</i> species, <i>Campylobacter</i> , <i>Aeromonas</i> , <i>Vibrio</i> species using Multiplex PCR. Each shipment has 4 simulated stool samples. Multiplex PCR panels and/or culture methods can be used. Participants are required to report the presence of one of the above pathogens or absence of the above pathogens
Covid 19 (SARS-CoV2)¹ 4 shipments/year	The program's objective is to provide EQA to those laboratories or testing facilities performing testing to detect the antigen or RNA of Covid 19 (Sars-Cov2). Each shipment consists of 4 samples.

* Procedure used to determine the Assigned Value:

¹ Assigned Values are determined by known values with results determined by specific proficiency test item formulation (e.g. manufacture or dilution).

² Assigned Values are determined by consensus values from expert participants.

³ Assigned Values are determined by consensus values from participants.

