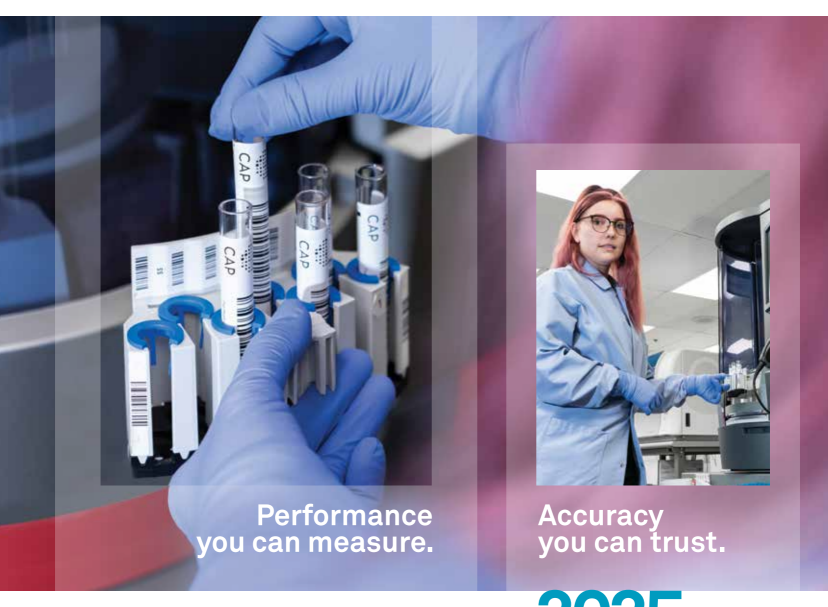


Surveys and Anatomic Pathology Education Programs



2025

Your Dedication, Our Gratitude— Supporting Patients' Every Step

A patient's journey can certainly feel daunting; it's riddled with unknowns at every turn, and often requires the efforts of an entire health care ecosystem. It takes so many distinct roles and responsibilities to ensure the best outcomes for patients, and some of those—by their very definition—tend to be more front and center than others.

But we feel quite comfortable saying that medical laboratory professionals are essential to the foundation of that ecosystem. Without you doing what you do, the whole journey could take a wrong turn. The accuracy of your work has a monumental impact on the overall health care system operating as smoothly as it should.

Moreover, we honor you and the care you take with every laboratory sample that helps shine a light on an otherwise intimidating path. We stand with you and the more than 23,000 laboratories worldwide that count on the CAP's PT/EQA programs to maintain high-quality, accurate patient testing.

The CAP also builds on its foundation of pathologist expertise by partnering with the 28 committees on our Council of Scientific Affairs to provide programs that promote excellence in the practice of pathology and laboratory medicine.

Together, let's continue our support for the patients and always strive for the best possible outcomes.

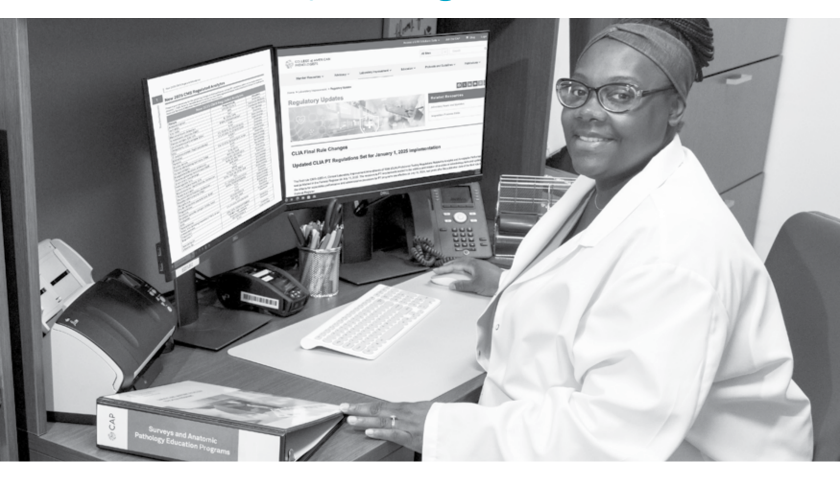


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Let us help you meet the new proficiency testing requirements.



You can count on the College of American Pathologists to guide you through CLIA's first major update to its proficiency testing requirements in more than 30 years.

- The CAP has reconfigured our PT/EQA programs where needed to meet the new participation requirements.
- For returning PT/EQA customers, we'll pre-populate your order forms during the order renewal process with the recommended programs based on your history.
- In our participant summaries, you'll find detailed information on the newly regulated analytes, revised grading criteria and acceptance limits, and all other critical changes.
- Your evaluation reports are always personalized with your laboratory's results, peer group statistics, and normalized results as the standard deviation index (SDI) to help you troubleshoot and identify bias, shifts, or trends.
- For CAP-accredited laboratories, PT enrollment will be audited in early 2025 to ensure your laboratory meets requirements.

See all the new CMS regulated analytes for 2025 on page 4.

Stay current with regulatory updates.



New Developments



We anticipate your needs because laboratory quality drives everything we do.

The CAP aligns to the updated CMS PT participation requirements, supporting laboratories that are subject to CLIA regulations and those that are CAP accredited.

Also new for 2025:

- Gastrointestinal panel created specifically for laboratories outside the US (GIPN)
- New program exclusively formulated to test proficiency of parathyroid hormone level (PTH)
- Linearity test covering a comprehensive range of thyroid analytes (LN50)

New Developments

New 2025 CMS Regulated Analytes	4
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New 2025 CMS Regulated Analytes

Prepare your laboratory for the upcoming changes in CLIA proficiency testing (PT) regulations, which will be implemented on January 1, 2025. Refer to the table below to review the new 2025 regulated analytes established by the Centers for Medicare & Medicaid Services (CMS) that meet all updated CLIA requirements.

New	2025 CMS Regulated An	alytes	
Analyte	Program Code	Discipline	Page
Acetaminophen	CZ/CZX/CZ2X, Z	Chemistry	54-56
Anti-HBs	VM1	Transfusion Medicine	243
Anti-HCV	VM1	Transfusion Medicine	243
Bacterial toxin detection	D, CDF5, GIP5	Microbiology	177, 187, 212
B-type natriuretic peptide (BNP)	BNP5, PCARM/PCARMX	Chemistry	59,64
Cancer antigen (CA) 125	K/KK	Endocrinology	82
Carcinoembryonic antigen (CEA)	K/KK	Endocrinology	82
Cholesterol, LDL	C1, C3/C3X, C4, CZ/CZX/CZ2X	Chemistry	54-56
CO ₂	C1, C3/C3X, C4, CZ/CZX/CZ2X	Chemistry	54-56
C-reactive protein (high-sensitivity)	HSCRP	Chemistry	63
Estradiol	Y/YY	Endocrinology	84
Ferritin	C1, C3/C3X, CZ/CZX/CZ2X, K/KK	Chemistry	54-56, 82
Folate, serum	K/KK	Endocrinology	82
Follicle stimulating hormone (FSH)	Y/YY	Endocrinology	84
Fungal antigen	F, F1, CRYP	Microbiology	194, 195
Fungal identification, molecular	MVP, VS, IDM5	Microbiology	191, 190, 209
Gamma glutamyl transferase (GGT)	C1, C3/C3X, CZ/CZX/CZ2X	Chemistry	54-56
Gram stain morphology	D, D2, D3, D5, RMC	Microbiology	177, 179, 180
Hemoglobin A1c	GH5, GH5I	Chemistry	62,63
Luteinizing hormone (LH)	Y/YY	Endocrinology	84
Mycobacterial identification, molecular	MTR5	Microbiology	193
Parasite antigen	P, P3, P4, P5, RML5	Microbiology	197, 198
Parasite identification, molecular	GIP5, TVG5, VS, STIM, MVP	Microbiology	212, 197, 190, 19
Parathyroid hormone (PTH)	PTH	Endocrinology	86
Phosphorus	C1, C3/C3X, CZ/CZX/CZ2X	Chemistry	54-56
Pro B-natriuretic peptide (pro-BNP)	BNP5, PCARM/PCARMX	Chemistry	59,64
Progesterone	Y/YY	Endocrinology	84
Prolactin	Y/YY	Endocrinology	84
Prostate specific antigen (PSA), total	K/KK	Endocrinology	82
Salicylate	CZ/CZX/CZ2X, Z	Chemistry	54-56
tCO ₂	AQ, AQH, AQIS	Blood Gas	92,93
Testosterone	Y/YY	Endocrinology	84
Total, iron-binding capacity (TIBC), direct measurement	C1, C3/C3X, CZ/CZX/CZ2X	Chemistry	54-56
Troponin I	CRT, CRTI, PCARM/PCARMX	Chemistry	60,64
Troponin T	NONE* High-sensitivity troponin T available	Chemistry	60
Vancomycin	CZ/CZX/CZ2X, Z	Chemistry	54-56
Vitamin B ₁₂	K/KK	Endocrinology	82

^{*} Conventional troponin T not available

2025 New Programs

	Quality Management Tools		
Subsection	Name	Program Code	Page
Quality Management Tools	Laboratory Staffing Ratios	QP251	25

General Chemistry and Therapeutic Drug Monitoring			
Subsection	Name	Program Code	Page
General Chemistry and Therapeutic Drug Monitoring	Waived Hemoglobin	HCC1	65

	Endocrinology		
Subsection	Name	Program Code	Page
Endocrinology	Parathyroid Hormone	PTH	86

Instrumentation Verification Tools			
Subsection	Name	Program Code	Page
Calibration Verification/Linearity	Thyroid Panel Calibration Verification/Linearity	LN50	136
Calibration Verification/Linearity	Factor VIII Calibration Verification/Linearity	LN51	133
Calibration Verification/Linearity	HBV Viral Load Calibration Verification/Linearity	LN52	133

Microbiology			
Subsection	Name	Program Code	Page
Parasitology	Trichomonas vaginalis, Molecular, 5 Challenge	TVG5	197
Parasitology	Rapid Malaria, 5 Challenge	RML5	198
Multidiscipline Microbiology	Gastrointestinal Panel, Global	GIPN	213

Transfusion Medicine, Viral Markers, and Parentage Testing				
Subsection	Name	Program Code	Page	
Transfusion Medicine	Transfusion Medicine With Electronic Crossmatch	JXM	232	
Transfusion Medicine	Transfusion Medicine—Automated With Electronic Crossmatch	JATXM	233	

Histocompatibility			
Subsection	Name	Program Code	Page
Histocompatibility	HLA Antibody Screen (Class I/Class II) Only	MXS	248
Histocompatibility	HLA Crossmatching, Antibody Screen, and Antibody Identification (Class I/Class II), Extra Plasma	MXEP	248

	Anatomic Pathology		
Subsection	Name	Program Code	Page
Surgical Pathology	CAP/NSH HistoQIP Pediatric Program	HQPED	288

2024 New Programs

Name	Program Code	Page
Quality Management Tools		
Technical Competency Assessment of Body Fluid Review for up to 25 Technologists,		
now called Assessment of Consistency of Body Fluid Morphologic Observations for up to 25 Technologists	QPB25	26
Quality Cross Check		
Quality Cross Check—High-Sensitivity Cardiac Markers	HCRQ	39
Quality Cross Check—Critical Care Blood Gas With Hematocrit	AQHQ	42
Quality Cross Check—Critical Care Blood Gas, i-STAT	AQSQ	42
General Chemistry and Therapeutic Drug Monitoring		
H. pylori Breath Test	НРВТ	75
Blood Gas, Critical Care, and Oximetry		
Critical Care Blood Gas With Hematocrit	AQH	92
Critical Care Blood Gas, i-STAT	AQIS	93
Instrumentation Verification Tools		
Cystatin C Calibration Verification/Linearity	LN49	135
Homotology and Clinical Microscopy		
Hematology and Clinical Microscopy	BCPV	1/0
Blood Cell Identification, Virtual	всгу	142
Microbiology		
Sexually Transmitted Infection Detection, Molecular	STIM	191
Mpox Molecular	MPOX	202
SARS-CoV-2 Molecular, 5 Challenge	COVM	203
SARS-CoV-2 Antigen, 5 Challenge	CVAG	203
Constice and Molecular Pathology		
Genetics and Molecular Pathology CAP/ACMG Applearniting Quantitation for Inheritad Metabolic Disorders	BGL4	250
CAP/ACMG Acylcarnitine Quantitation for Inherited Metabolic Disorders	DUL4	259
Anatomic Pathology		
HER2 and ER Immunohistochemistry Interpretation Only	HERI	298
Navigating Multimodality Biomarker Assessment	NMBA/NMB1	300

Continuing Education



We support laboratory professionals. Maintain your certification with Surveys continuing education (CE).

- Offer your staff more than 100 CE credits.
- Advance skills with education activities developed by more than 600 physicians and doctoral scientists with expertise in pathology and laboratory medicine.
- Meet certification and licensure requirements with CE across multiple disciplines.

Continuing Education

Continuing Education Programs	. 8
Competency Assessment Hub	15
QMEd™ Online Educational Courses	18

Discontinued Programs

Informatics Essentials for Pathologists (ICBE/ICBE1)

Continuing Education Programs

Your laboratory demonstrates its commitment to quality by choosing CAP Surveys programs. You'll find the same level of quality in the CAP Continuing Education Programs.



CME (Continuing Medical Education for Physicians)

Accreditation

The College of American Pathologists (CAP) is accredited by the Accreditation Council for Continuing Medical Education (ACCME) to provide continuing medical education for physicians.

CME Category 1

The CAP designates these educational activities for a maximum of the stated number of *AMA PRA Category 1 Credits*™. Physicians should claim only the credit commensurate with the extent of their participation in the activity.



credif CE (Continuing Education for Nonphysicians)

The CAP designates these educational activities for a maximum of the stated number of credits of continuing education. Participants should claim only the credit commensurate with the extent of their participation in the activity.

The American Society for Clinical Pathology (ASCP) Board of Certification (BOC) Certification Maintenance Program (CMP) accepts these activities to meet its continuing education requirements.

These activities are approved for continuing education credit in California and Florida.

Cytotechnologists may apply the credits from the PAP Education (PAPCE/PAPJE/PAPKE/PAPLE/PAPME), NGC, FNAG, FNA, and TICP programs toward the required educational activities for the American Society of Cytopathology (ASC) Continuing Education Credit Program (CECC) and the International Academy of Cytology (IAC).



credit This activity is eligible for continuing medical education (CME) or continuing education (CE) credit.

Surveys Continuing Education Activities

When your laboratory participates in CAP Surveys, every member of your team can enroll in education activities and earn continuing education (CE) credit at no additional charge. Simply follow these steps:

- 1. Establish a free online account.
- 2. Complete a reading provided in the Participant Summary or Final Critique.
- 3. Answer online learning assessment questions.
- 4. Claim CE certificate.

Each member of your staff can access the Surveys education activities for a maximum of 12 months.

Surveys Educational Activities								
Program Name	Program Code	Discipline	Catalog Page(s					
General Chemistry	C1, C3/C3X, C4, CZ/CZX/CZ2X, Z	Chemistry	54-56					
Blood Gas	AQ, AQH, AQIS	Chemistry	92-93					
Endocrinology	K/KK	Chemistry	82					
Sex Hormones	Y/YY	Chemistry	84					
Quality Cross Check—Whole Blood Glucose	WBGQ	Chemistry/Quality Cross Check	37					
Coagulation—Limited	CGB, CGDF, CGL	Coagulation	166					
Blood Cell Identification, Photographs Blood Cell Identification, Virtual	BCP, BCPV	Hematology and Clinical Microscopy	142					
Bone Marrow Cell Differential	BMD	Hematology and Clinical Microscopy	145					
Hematology Automated Differential Series	FH1-FH4, FH9-FH10, FH13, FH16-FH17	Hematology and Clinical Microscopy	140					
Hematology—Basic	HE	Hematology and Clinical Microscopy	140					
Virtual Body Fluid	VBF	Hematology and Clinical Microscopy	154					
Immunology	ANA, ASO, CRP, HCG, IM, RF/RFX, RUB/RUBX, IL, IG/IGX, S2, S4, S5, AHT, CCP, RDS, G	Immunology and Flow Cytometry	216, 217 218, 220 222					
Special Chemistry	M, OLI, SPE, UBJP	Chemistry	74, 76					
Bacteriology	D	Microbiology	177					
Mycobacteriology	E	Microbiology	193					
Mycology and Aerobic Actinomycetes	F	Microbiology	194					
Yeast Identification	F1	Microbiology	194					
Limited Bacteriology	D1, D2, D3, D5, D6, D8, MC3, MC4, RMC	Microbiology	179-180 182-183					
Embryology	EMB	Reproductive Medicine	163					
Sperm Count, Motility, Morphology, and Viability	SMCD, SM1CD, SM2CD	Reproductive Medicine	162					
Semen Analysis	ASA, SC, SC1, PV, PV1, SM, SV	Reproductive Medicine	162					
Toxicology	DFC, NOB, OFD, SCDD, VF	Toxicology	108, 111 103, 104					
Transfusion Medicine	J, JXM, JE1, JAT, JATXM, JATE1, J1	Transfusion Medicine	232-233					

Surveys Self-Reported Training Opportunities

When your laboratory participates in CAP Surveys, every member of your team can receive self-reported training opportunities.

Self-Repo	orted Training Oppo	rtunities*	
Program Name	Program Code	Source	Catalog Page(s)
Quality Management Tools		'	
Laboratory Staffing Ratios	QP251	Data Analysis and Critique	25
Assessment of Consistency of Body Fluid Morphologic Observations	QPB10, QPB25	Data Analysis and Critique	26
Assessment of Consistency of Peripheral Blood Morphologic Observations	QPC10, QPC25	Data Analysis and Critique	27
Assessment of Consistency of Gram Stain Morphologic Observations	QPD10, QPD25	Data Analysis and Critique	28
Hematology and Clinical Microscopy			
Blood Cell Identification, Photographs/Virtual	BCP, BCPV	Participant Summary	142
Bone Marrow Cell Differential	BMD	Participant Summary	145
Expanded Virtual Peripheral Blood Smear	EHE1	Participant Summary	150
Hematology Automated Differential Series	FH1-FH4, FH9-FH10, FH13, FH16-FH17	Participant Summary	140
Hematology—Basic	HE	Participant Summary	140
Hemoglobinopathy	HG	Participant Summary	147
Virtual Body Fluid	VBF	Participant Summary	154
Virtual Peripheral Blood Smear	VPBS	Participant Summary	149
Clinical Microscopy	CMP, CMMP, CMP1	Participant Summary	152-153
Microbiology			
Blood Parasite	BP	Participant Summary/Final Critique	198
Expanded Bacteriology	DEX	Participant Summary/Final Critique	178
Yeast	F1	Participant Summary/Final Critique	194
Parasitology	Р	Participant Summary/Final Critique	197
Ticks, Mites, and Other Arthropods	TMO	Participant Summary	198
Worm Identification	WID	Participant Summary	198
Toxicology			
Drug Monitoring for Pain Management	DMPM	Participant Summary	110
Drug Monitoring for Pain Management	DMPM	Participant Summary	

^{*}Notes:

[•] CAP Self-Reported Training Opportunities do not offer CE credit, but can be used toward fulfilling requirements for certification maintenance by agencies such as the American Society for Clinical Pathology (ASCP). Please verify with your certifying agency to determine your education requirements.

[•] These opportunities are subject to change. Refer to the Participant Summary/Final Critique for availability.

Continuing Certification (CC)

Continuing Certification (CC) is the board certification program that involves continuous professional development and ensures that an American Board of Pathology (ABPath) board-certified pathologist is committed to lifelong learning and competency in a specialty and/or subspecialty.

There are six competency categories defined by the American Board of Medical Specialties (ABMS) and endorsed by the ABPath to fulfill specific CC requirements. They are listed below with their descriptions.

All CAP education activities providing CME credits meet the CC Part II: Lifelong Learning requirements. Some programs will meet the requirements for CC Improvement in Health and Health Care (IHHC) (formerly Part IV) at the laboratory or the individual level. Programs that meet IHHC are identified within the description of the program.

Interpersonal and Communication Skills

Demonstrate interpersonal and communication skills that result in effective information exchange and teaming with patients, patients' families, and professional associates.

Medical Knowledge

Demonstrate knowledge of established and evolving biomedical, clinical, and cognate sciences and the application of this knowledge to pathology.

Practice-Based Learning and Improvement

Demonstrate ability to investigate and evaluate diagnostic and laboratory practices in your own laboratory, to appraise and assimilate scientific evidence, and to improve laboratory practices and patient care.

Patient Care

Demonstrate a satisfactory level of diagnostic competence and provide appropriate and effective consultation in the context of pathology services.

Professionalism

Demonstrate a commitment to carrying out professional responsibilities, adherence to ethical principles, and sensitivity to diverse patient population.

Systems-Based Practice

Demonstrate understanding of and contribution to local, regional, and national health care systems, and support health care in systems-based practice definition.

Education Programs									
Program Name	Program Code	Maximum AMA PRA CME Category 1 Credits Annually	Maximum CE Credits Annually	Format	Catalog Page				
Autopsy Pathology*	AUP/AUP1	12.5	12.5	Online (DigitalScope®)	302				
Clinical Pathology Improvement Program*	CPIP/CPIP1	15	NA	Online	14				
Digital Slide Program— Dermatopathology*	DPATH/DPATH1	15	NA	Online (DigitalScope)	303				
Digital Slide Program in FNA*	FNA/FNA1	10	10	Online (DigitalScope)	311				
Fine-Needle Aspiration Glass Slide	FNAG/FNAG1	10	10	Glass Slides	312				
Forensic Pathology*	FR/FR1	12.5	12.5	Online	314				
Hematopathology Online Education*	HPATH/HPATH1	12.5	12.5	Online (DigitalScope)	151				
Nongynecologic Cytopathology Education**	NGC/NGC1	25	25	Glass Slides With Online Cases (DigitalScope)	310				
Navigating Multimodality Biomarker Assessment*	NMBA/NMB1	4	4	Online (DigitalScope)	300				
Neuropathology Program*	NP/NP1	10	NA	Online (DigitalScope)	305				
Gynecologic Cytopathology PAP Education Program***	PAPCE/APAPCE PAPJE/APAPJE PAPKE/APAPKE PAPLE/APAPLE PAPME/APAPME Series 1 or 2	/APAPCE /APAPJE /APAPKE /APAPLE /APAPME		Glass Slides	307				
Glass Slide Cytopathology PAP PT Program (With Glass Slide PAP Education)***	PAPCPT/APAPCPT PAPJPT/APAPJPT PAPKPT/APAPKPT PAPLPT/APAPLPT PAPMPT/APAPMPT	8	8	Glass Slides	306				
Performance Improvement Program in Surgical Pathology	PIP/PIP1	40	NA	Glass Slides With Online Cases (DigitalScope)	283				
Online Performance Improvement Program in Surgical Pathology*	PIPW/PIPW1	40	NA	Online (DigitalScope)	282				
Nongynecologic Cytopathology Intraoperative Touch Imprint/ Crush Preparation Program*	TICP/TICP1	10	10	Online (DigitalScope)	309				
Virtual Biopsy Program*	VBP/VBP1	25	NA	Online (DigitalScope)	284				

^{*}Program is available for purchase online. Go to cap.org and choose the Education tab.

System Requirements

DigitalScope is a web-based whole slide image (WSI) retrieval and viewing system. **The current version, DSv6.0, does not require Microsoft Silverlight.** DigitalScope is supported by the latest Chrome and Firefox releases, and the last two major Edge and Safari versions.

Find current information on system requirements on cap.org; click **Browser and Operating System Requirements** at the bottom of the homepage. Download speeds and appearance will vary depending on your internet connection, browser, and computer power.

^{**}NGC provides up to 20 CME/CE credits for the glass slides and five CME/CE credits for the online slide portion of the program.

^{***}PAP provides up to eight CME/CE credits for the glass slides.

Navigating Multimodality Biomarker Assessment NMBA/NMB1						
Program Name	Program Code Cases per Mailing					
Multimodality biomarker assessment case analysis		ı	2			

Biomarkers can be tested through a variety of methodologies (eg, flow cytometry, immunohistochemistry, next-generation sequencing, PCR-based testing, karyotype, FISH, clinical chemistry). Each modality has different technical strengths and weaknesses, along with varied analytical and clinical specifications such as sensitivity and specificity. Results may not be clearly concordant across the modalities. This program will challenge pathologists to resolve discrepancies between different methods or "multimodality" biomarker testing.

Program Information

- NMBA Online program, whole slide images powered by DigitalScope technology (if available); NMBA provides CME or CE for one pathologist or laboratory professional
- NMB1 Reporting option with CME or CE credit for each additional pathologist or laboratory professional (within the same institution); must order in conjunction with program NMBA
- Two mailings per year with two cases each mailing
- Earn a maximum of four CME credits (AMA PRA Category 1 Credits™) per pathologist and a maximum of four CE credits per laboratory professional per year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHCC).
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



Access CPIP cases when and where it's convenient via PC or personal mobile device.

Pathologists can keep abreast of current scientific knowledge with interactive, case-based learning addressing common issues faced in the laboratory.

CPIP supports clinical pathologists as well as anatomic pathologists who cover clinical pathology. A diverse portfolio of real-life case scenarios, including images and clinical background, helps pathologists to stay current on issues and advances in the laboratory.

CPIP is designed for pathologists, by pathologists. Each case is developed and peer-reviewed, ensuring learning is practical and easily applied to work. Thought-provoking questions with feedback and multiple choice knowledge checks assess and confirm diagnostic skills. Participants may apply 1.25 CME credits for each CPIP toward the ABPath's Continuing Certification (CC) requirements.

Clinical Pathology Improvement Program CPIP/CPIP1						
Program Name Program Code Cases per Year						
Online cases in clinical pathology	ı	12				

Consider CPIP for:

- Medical directors seeking to continuously improve the collective skills and clinical pathology knowledge of their team
- · Pathologists with clinical and/or laboratory management responsibilities
- · Pathologists seeking CME CC credits in clinical pathology
- · Subspecialty clinical pathologists who need to keep current

Discipline	Case Schedule (subject to change)	Month 2025
Laboratory Management	Provider Performed Microscopy (PPM) Testing	January
Transfusion Medicine	ABO Discrepancies	February
Transfusion Medicine	Blood Bank Regulations	March
Chemistry	Hyperbilirubinemia	April
Hematology	Hematologic Pleural Effusions	May
Microbiology	Bloodstream Infections	June
Laboratory Management	CLIA Director Responsibilities and Risks	July
Cytogenetics	Plasma Cell Myeloma	August
Transfusion Medicine	Transfusion Reactions	September
Molecular Pathology	Pharmacogenomics	October
Hematology	Peripheral Blood Smear - Part 1	November
Hematology	Peripheral Blood Smear - Part 2	December

To learn more visit cap.org and search CPIP.

Program Information

- CPIP One online clinical laboratory case per month
- CPIP1 Additional pathologist (within the same institution) reporting option with CME credit; must order in conjunction with CPIP
- Earn a maximum of 15 CME credits (AMA PRA Category 1 Credits™) per year.
- Twelve cases per year; your CAP shipping contact will be notified via email when the activity is available



Competency Assessment Hub

A single platform for maintaining your staff competency records and providing CE credits

Presented in partnership with MediaLab, the CAP Competency Assessment Hub helps individual laboratories and entire health care networks ensure they meet CLIA competency assessment requirements and fulfill laboratory professional continuing education (CE) needs. Built on MediaLab's platform, the CAP's Competency Assessment Hub helps you stay in compliance and avoid being cited for a deficiency by managing your personnel's training and competency assessment performance and records.

- System/network subscriptions now available—Enroll your entire system and participate as individual, linked sites. Standardize your competency assessments across the system and provide centralized documentation.
- Customizing tools—The question bank lets you design your own assessment courses to match your laboratory's written procedures. Checklist tool, CourseBuilder, and Compass competency assessments can ensure convenient documentation for all six areas of competency as defined by CLIA and the CAP Laboratory Accreditation Program.
- Auto-assignment of assessments and reminder emails—Never forget your staffs' next assessments.
- Intuitive reporting—With just a few clicks, administrators can stay on top of documentation and records to track progress toward required dates and training for all staff members.
- Instrument-specific checklists—More than 130 standard checklists help you meet your laboratory's documentation needs.
- High-quality Pro courses—Your laboratory staff can earn PACE CE credits in a variety of disciplines and courses.
- Easy online access—The Competency Assessment Hub is cloud-based, so it's available 24/7 from any PC, laptop, or tablet—wherever you have an internet connection.

Add Safety & Compliance Courses especially developed for the laboratory

As an add-on option, the Competency Assessment Hub offers a package of nine complementary safety and compliance courses with PACE CE credits. The package is appropriate for annual laboratory-specific compliance training and for clinical laboratory science students prior to clinical rotations. These courses include:

- · OSHA Bloodborne Pathogens
- · OSHA Hazard Communication and Chemical Hygiene
- OSHA Electrical Safety
- OSHA Fire Safety
- · OSHA Formaldehyde
- Tuberculosis Awareness for Healthcare Workers
- Medical Error Prevention: Patient Safety
- · Ethics and Code of Conduct in Healthcare
- · HIPAA Privacy and Security Rules

With the Competency Assessment Hub, you can keep your laboratory and network organized and inspection-ready every day of the year. Choose the Competency Assessment Hub subscription that best fits your needs. Please refer to the ordering information and course descriptions on the following pages. For more information, visit cap.org and choose Competency Assessment Hub from the Education Main Page via the Education tab.

Number of Users*	Competency Assessment Hub	Competency Assessment Hub With Optional Safety & Compliance Courses**
2 to 50	CA0050	CA0050 + XCA0050
51 to 250	CA0250	CA0250 + XCA0250

^{*}For subscriptions for single users or more than 250 users and networks, please contact the CAP for more information.

^{**}Safety & Compliance Course subscriptions require a standard Competency Assessment Hub subscription.

2025 Pro Courses

Blood Bank/Transfusion Medicine

- · ABO typing discrepancies
- · Antibody screen and identification
- Direct antiglobulin test
- Blood components—storage, handling, and selection
- · Transfusion reactions
- Quality control in the blood bank laboratory

Chemistry

- · Cardiac biomarkers
- · Liver and renal testing
- · Electrolytes, acid base, and anion gap
- Clinical toxicology
- · Therapeutic drug monitoring
- Chemistry QC, calibration, and reportable range

Hematology/Coagulation

- · Erythrocyte morphology
- · Erythrocyte inclusions
- · White blood cells
- · White blood cell inclusions
- Common coagulation tests
- Platelet testing, morphology, and disorders

Histology

- Immunohistochemistry—part 1
- Immunohistochemistry—part 2
- · Special stains
- · Histology specimen handling
- · Quality management in histology
- Safety issues in the histology laboratory

<u>Immunology</u>

- · Hepatitis testing
- · Qualitative HIV testing
- Human chorionic gonadotropin and fetal fibronectin
- · Rapid serology kit tests
- Molecular amplification methods for detection of infectious diseases
- Monitoring the testing process in immunology

Microbiology

- Gram stain: organism detection and differentiation
- · Urine and body fluid cultures
- · Genital tract pathogens
- · Blood cultures
- Microbiology of the gastrointestinal tract
- · The microbiology of wounds

Phlebotomy/Specimen Processing

- · Venipuncture
- Challenges of phlebotomy: pediatric blood collection, alternate sites, and difficult draws
- Phlebotomy professionalism and ethics
- Common pitfalls in specimen processing
- Specimen collection for workplace urine drug testing programs and forensic drug and alcohol testing
- General specimen handling and transportation requirements

Point-of-Care Testing



- · Urine dipstick
- Whole blood prothrombin time and INR (PT/INR) testing
- · Whole blood glucose testing
- · Cardiac biomarkers
- · Blood gas testing
- Provider-performed microscopy and limited waived testing

Quality Programs/Management

- · New instrument method validation
- · Monitoring the quality control program
- · Document control
- Investigating occurrences (occurrence reports, root cause analysis, and corrective action)
- · Competency evaluation
- Development and implementation of a quality management program

Safety

- · General laboratory safety
- · Bloodborne pathogens
- Laboratory waste and spill management
- · Fire and electrical safety
- · Hazardous chemicals
- SARS-CoV-2/COVID: biosafety precautions
- · Ergonomics

Urinalysis/Body Fluids

- · Physical and chemical urinalysis
- Microscopic urinalysis—part 1
- Microscopic urinalysis—part 2, crystals and casts
- · Cerebrospinal fluid analysis
- · Serous and synovial fluids
- · Semen analysis

Safety & Compliance Courses

OSHA Bloodborne Pathogens—Addresses the OSHA Bloodborne Pathogens standard as it applies to clinical and medical laboratories. Covers major bloodborne pathogens, including hepatitis B and HIV. Focuses on proper handling of sharps, personal protective equipment (PPE), engineering controls such as microbiological safety cabinets, and proper work practices like handwashing.



OSHA Hazard Communication and Chemical Hygiene—Describes the OSHA Chemical Hygiene Standard and helps satisfy OSHA requirements for annual training. Explains Haz-Com, the National Fire Protection Agency diamond, the Safety Data Sheet, and common-sense laboratory safety rules applied to clinical laboratory practice.

OSHA Electrical Safety—Addresses electrical safety and electrical hazards commonly found in the clinical laboratory. Covers prevention and safety measures, fighting electrical fires, and treatment of electrical injuries.

OSHA Fire Safety—Teaches the basics of fire safety in the clinical laboratory, including classes of fire and key acronyms, such as PASS and RACE. Addresses fire prevention, drills, and firefighting techniques.

OSHA Formaldehyde—Covers essentials for any laboratory that uses formaldehyde or formalin. Shares facts about formaldehyde, safety risks, proper handling procedure, monitoring, spill cleanup, and PPE.

Tuberculosis Awareness for Health Care Workers—Provides background information about the spread of tuberculosis, purified protein derivative (PPD) testing procedures, CDC guidelines, and methods of control.

Medical Error Prevention: Patient Safety—Includes potential causes of medical errors in the clinical laboratory, important legislation and definitions, and steps laboratory professionals can take to reduce the impact of medical errors in their workplace. Serves as an ideal part of an effective medical error reduction program. Appropriate for both experienced and newer laboratory personnel.

Ethics and Code of Conduct in Health Care—Designed to guide health care employees on the importance of ethics and code of conduct by outlining privacy and patient health information regulations, conflict of interest, professional competence, effective communication, and more.

HIPAA Privacy and Security Rules—Addresses the Health Insurance Portability and Accountability Act of 1996 (HIPAA) privacy regulations and treatment of protected health information (PHI) in a succinct manner. Content is directed at laboratory staff, from desk personnel to phlebotomists to medical technologists. Includes technical and physical safeguards, minimum necessary standards, administrative requirements, and authorization.

Identify and control risks in your laboratory.

The Risk Management QMEd online course provides a realistic case study as well as video commentary by CAP pathologists, inspectors, and ISO 15189 assessors. Learn how to:

- Find, prioritize, and control risks
- Use common tools
- Assess how your laboratory's culture is affecting risks

Includes a Risk Register Tool spreadsheet, which helps prioritize and keep track of risks.

See the Continuing Education section. Add QMEDRISK to your order. "Managing risks is a mindset that needs to be present throughout the laboratory ... This course will help you manage risk to a level that is acceptable to our physicians, our patients, and our administration."

> Dr. Gaurav Sharma, MD, FCAP Division Head of Regional Laboratories Henry Ford Health System

QMEd™ Online Educational Courses

Tailored education and quality tools developed with pathologist input



Quality Management Educational Resources (QMEd) courses will help you:

- Build a quality management system (QMS)—one piece at a time—that sustains your continuous improvement and Lean efforts.
- Self-assess your current QMS against international quality standards.
- · Interpret ISO 15189 requirements.
- · Perform internal audits using tracer audit and process audit methods.
- Implement and refine occurrence management with root cause analysis.

Course Information

- · Delivered on demand so you can pause, resume where you left off, and learn at your own pace
- · Mobile-friendly so that you can learn where and when you want
- · Accessible for a minimum of twelve months
- Includes continuing education (CE) credit
- · Individual learners use their own login with the ability to bookmark the course and continue where they left off.

About the Courses

Risk Management Order QMEDRISK

Learn how different elements of the quality management system—internal audit, data analysis, etc—play a role in identifying and controlling risk. Learn best practices for managing risk, plus practical tools for all phases of the risk management process. Includes a case example showing how high-level risk assessment can be integrated into management review.

4 CE credits available

Quality Culture Order QMEDOCUL

This program—designed for laboratory medical directors, administrative directors, quality managers, and other leaders whose decisions affect the culture of their laboratory—provides an adaptable program for proactive culture change. Its unique Culture Assessment Tool helps laboratory leadership get a picture of where your organization is strong and where it needs to improve, then helps make culture change a reality. It also includes video commentary by CAP member pathologists.

4 CE credits available

Root Cause Analysis Order QMEDROOT

Designed for laboratory quality managers and implementation team members. Learn real-world methodology and tools to conduct and implement a root cause analysis, performing key steps based on a participant case study. Choose further examples based on your work setting (eg, hospital, reference laboratory, or contract research organization). Includes the RCA Performance and Feedback Toolkit, which an organization can use to guide and assess root cause analysis projects.

6 CE credits available

Mistake Proofing Order QMEDMIST

Learn to develop and revise processes, reduce errors, and handle risks. The course methodology is focused on five main categories of mistake-proofing tactics, with examples taken from laboratory medicine. It includes video commentary by CAP member pathologists who have experience using Lean and other process improvement techniques.

4 CE credits available

Internal Auditing Order QMEDAUDT

Improve your internal audit capability with a proven methodology for process, tracer, and laser audits. Learn to prepare for interviews, communicate findings to your quality management team, and use audits to drive process improvements. Includes detailed, real-world examples you can use to build your own audit plans, plus multimedia presentations of key concepts.

3 CE credits available

Management Review Order QMEDMGMT

Understand the ISO 15189 requirements for management review. CAP ISO 15189 assessors cover structuring review meetings, communicating results, and prompting strategic management decisions—all to benefit your organization's health.

2 CE credits available

Quality Manual Development Order QMEDMANL

Go beyond a quality plan—develop a manual that organizes and communicates your laboratory's quality management system. The course materials include a well-written and effective sample manual, which you can use to organize and create your own. Plus, the CAP's ISO 15189 assessors demonstrate how to link your quality policy to quality objectives and metrics.

2 CE credits available

Document Control Order QMEDDOCU

This "how-to" course details how to control documents to meet ISO 15189 requirements, how to accomplish document control even with minimal resources (such as spreadsheets), and how document control contributes to cost containment. The CAP's ISO 15189 assessors provide commentary on common pitfalls and best practices.

2 CE credits available

QMS Implementation Roadmap Order QMEDROAD

Outlines the practical steps necessary to build, implement, and maintain a quality management system that meets the ISO 15189 standard. Video recordings of the CAP's ISO 15189 assessors provide perspective on best practices and pitfalls. Designed for laboratory quality managers and implementation team members.

2 CE credits available

15189 Walkthrough Order QMEDWALK

This course summarizes each main clause of the ISO 15189 standard, clarifying its intent and key requirements. CAP assessors offer context in videos that also provide examples of how technical problems relate to fundamental deficiencies in the quality management system. This course, designed for laboratories considering implementation, is updated for the ISO 15189:2022 edition.

2 CE credits available

Make sure your laboratory team is ready to meet the challenges ahead. Add QMEd courses to your order form. For more information, visit cap.org and search QMEd.

Expand your expertise with Root Cause Analysis.

Developed with pathologist input, the Root Cause Analysis QMEd online course is infused with real-world laboratory examples, giving you confidence in:

- Using root cause analysis tools
- Recognizing common pitfalls
- Performing key steps Applying best practices

You'll receive our unique Root Cause Analysis Toolkit to help communicate best practices and provide feedback to project teams, with the goal of solving problems permanently.

See the Continuing Education section. Add QMEDROOT to your order.

"WOW! Very impressive training module. Probably the best self-taught module I've seen in years. Very systematic, very visual, very easy to follow ... staying with the tried and true textbook of **Root Cause Analysis."**

> Jim Ellis Managing Partner MME Consulting, LLC

Take your quality system to the next level.

The CAP 15189SM Accreditation Program provides accreditation to the ISO 15189:2022 4th edition, an international standard to recognize quality and competence in medical laboratories.

Our program offers:

- A distinct approach, using the CAP Laboratory Accreditation Program as a foundation
- Dedicated, expert assessors who specialize in ISO 15189
- Unique, tailored education and quality tools developed with pathologist input
- A personalized, flexible accreditation process

Contact us to learn more at cap15189@cap.org.



Quality Management Tools



Easily integrate quality improvement into your daily work processes.

Measure and document your process improvements with these convenient tools:

- Assess clinical laboratory staffing ratios and benchmark performance (QP251).
- Streamline your efforts to assess consistency of morphologic observations of peripheral blood (QPC10/ QPC25), Gram stains (QPD10/QPD25), and body fluid review (QPB10/QPB25).

Quality Management Tools

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Quality Management Tools

Benchmark outside your laboratory.

The CAP Quality Management Tools can improve your total testing process by providing a convenient solution to measure and document process improvements within your laboratory's quality management system.

- Short-Term Quality Studies and Morphology/Competency Assessments provide opportunities to benchmark performance indicators, compare normative rates, and assist your laboratory in meeting checklist requirements.
- Continuous Quality Monitors examine performance indicators such as turnaround time and specimen identification errors throughout the year.

Available for clinical pathology laboratories, Quality Management Tools examine preanalytic, analytic, and postanalytic phases, helping participants to:

- Establish realistic goals by comparing their performance against other institutions with comparable demographics.
- Monitor progress through unique and robust quality indicators on a periodic basis.
- Make effective decisions based on practical and in-depth quality management reports.
- Improve efficiencies to allow time for more patient-centric activities.
- Easily integrate quality improvement into their daily work processes.
- **Meet requirements** of the CAP Laboratory Accreditation Program checklists and The Joint Commission standards.

Purchase combination packages and save.

2025 Short-Term Quality Studies and Morphology/Competency Assessments						
Module/Package	Program Code					
Individual Short-Term Quality Studies and Morphology/Competency Assessments	QP251, QPB10, QPB25, QPC10, QPC25, QPD10, QPD25					
Four Quality Management Tools (QP251, QPB10, QPC10, QPD10)	PRO					

2025 Continuous Quality Monitors						
Module/Package	Program Code					
Individual Continuous Quality Monitors	QT2, QT3, QT4, QT7, QT8, QT10, QT16, QT17					
Clinical Pathology Module—includes all eight Continuous Quality Monitors	QTC					

Complement your quality management program needs with these clinical pathology studies.

Clinical Pathology Study	Testing Phase Purpose										
Select from the following studies to support your quality improvement initiatives.	Preanalytic	Analytic	Postanalytic	Anatomic Pathology	Clinical Pathology	Turnaround Time	Patient Safety	Microbiology	Transfusion Medicine	Chemistry/ Hematology	Customer Satisfaction
Laboratory Staffing Ratios QP251 (QPR-A) NEW	•	ı		1			ı	ı			
Assessment of Consistency of Body Fluid Morphologic Observations (QPB10/QPB25)		ı	ı		ı		ı			ı	
Assessment of Consistency of Peripheral Blood Morphologic Observations (QPC10/QPC25)		ı	ı		ı		ı			I	
Assessment of Consistency of Gram Stain Morphologic Observations (QPD10/QPD25)		ı	ı		ı		ı	ı			
Blood Culture Contamination (QT2)	ı	ı					I	ı			
Laboratory Specimen Acceptability (QT3)	ı										I
In-Date Blood Product Wastage (QT4)			ı		ı		ı				
Satisfaction with Outpatient Specimen Collection (QT7)							ı				
Stat Test Turnaround Time Outliers (QT8)		ı			ı		ı				
Critical Values Reporting (QT10)			ı				ı				
Corrected Results (QT16)			ı				ı	ı			
Outpatient Order Entry Errors (QT17)	ı						I	ı			

The CAP requires accredited laboratories to have a quality management plan that covers all areas of the laboratory and includes benchmarking key measures of laboratory performance (GEN.13806, GEN.20316, COM.04000). The Joint Commission requires accredited hospitals, laboratory staff, and leaders to regularly collect and analyze performance data (PI.01.01.01, Pl.03.01.01, LD.03.06.01, LD.03.07.01). CLIA requires laboratories to monitor, assess, and correct problems identified in preanalytic, analytic, and postanalytic systems (§493.1249, §493.1289, §493.1299).

Short-Term Quality Studies and Morphology/Competency Assessments

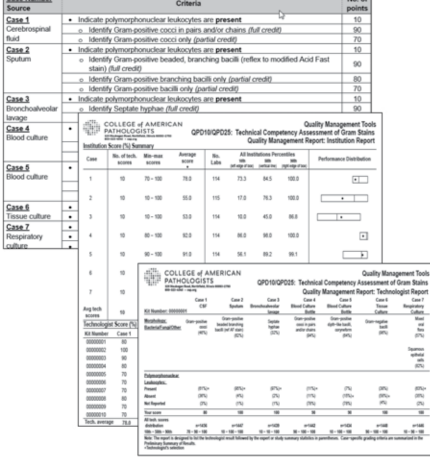
Implement quality monitoring—Use these comprehensive short-term quality studies and morphology/competency assessments to learn how to start monitoring and measuring key processes that may not be commonly monitored in your laboratory. These assessments also analyze emerging industry trends and topics to keep your laboratory ahead of the curve.

Gain experience in data collection and analysis—Based on data collected and submitted between predetermined dates, the CAP provides personalized reports with the individual participant's performance compared against peers.

Strengthen your quality assessment expertise—CAP pathologist experts provide in-depth discussions and identify best practices for laboratories to strive for. In addition, the studies' consolidated results are carefully reviewed and analyzed to be published in the form of scientific articles for further analysis.

Participating laboratories receive:

- · User Guides
- Templates and instructions for data collection
- Individual Participant Summaries and interpretation guides
- Data distributions and initial analysis of laboratory practices
- Data Analysis and Critiques with author commentaries on improvement opportunities, dependent on study type and complexity
- Morphology/Competency Program inclusions:
 - o Participant Summary Reports have alllaboratories, institution, and individual study results.
 - o Data Analysis and Critique has commentaries from subject matter experts on the importance of each case with links for viewing.



Laboratory Staffing Ratios QP251 (QPR-A)



Introduction

Laboratory staff play an important role in the detection, diagnosis, and treatment of disease by performing tests in laboratories. These staff account for two-thirds of direct clinical laboratory costs. This QP study is designed to produce data that will assist laboratory managers and directors in managing those costs and gauging their staffing levels.

Laboratories participating in this study will submit data on their staffing levels for laboratory sections, and may participate in any or all areas including anatomic pathology, chemistry/hematology/immunology, microbiology, molecular pathology, phlebotomy, point-of-care testing, and transfusion medicine. From these levels, staffing ratios will be calculated for these sections relative to managerial staffing and billable tests. Each laboratory's staffing ratios will be compared against those of other institutions participating in this study, and where applicable, against peer groups with similar billable test profiles.

Enrollment in this study will help laboratory directors address CAP Laboratory Accreditation Program Checklist statement DRA.11300, which requires sufficient numbers of personnel to be available to meet the needs of the laboratory, and The Joint Commission Standard HR.01.02.05, which requires the laboratory to have the necessary staff to support the services it provides.

Objectives

The aims of this study are to measure staffing levels in different areas of the laboratory, calculate key staffing ratios and levels, and compare all staffing ratios with those of other institutions participating in this study.

Data Collection

Participants will use their laboratory's or institution's revenue and usage reports to obtain billable test counts and staffing figures for the most recently completed fiscal year.

Performance Indicators

- · Anatomic Pathology
 - O Histology blocks/Histology non-management FTE
 - o Cytology accessions/Cytology non-management FTE
 - O Non-management FTE/Management FTE
- Chemistry/Hematology/Immunology
 - o Total billable tests/Non-management FTE
 - o Non-management FTE/Management FTE
- Microbiology
 - o Total billable tests/Non-management FTE
 - O Non-management FTE/Management FTE
- · Molecular Pathology
 - o Total billable tests/Non-management FTE
 - O Non-management FTE/Management FTE

Phlebotomy

- Total inpatient blood draws/Inpatient phlebotomist FTE
 Total outpatient blood draws/Outpatient phlebotomist FTE
- Point-of-Care Testing (POCT)
 - o POCT billable tests/Laboratory FTE overseeing POCT
- · Transfusion Medicine
 - o Crossmatches or type and screens/Non-management FTE
 - o Transfused units/Non-management FTE
 - O Non-management FTE/Management FTE

This is a one-time study conducted in the first quarter.

Assessment of Consistency of Body Fluid Morphologic Observations QPB10/QPB25

Introduction

Laboratories receive a variety of body fluids for evaluation that technologists review. Technical staff must maintain their identification skills of these specimens, and laboratories are required to provide education and to assess consistency of reporting morphology among staff and competency of body fluid cell identification on an annual basis.

Objectives

This study will assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the performance of accurate body fluid cell counts and identification of other body fluid features. Results of this study will assist individuals, the laboratory director, and the manager with areas to focus on for improvement and education.

The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for consistency of reporting morphology among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

Technologists will access a series of online, whole slide images to assess their ability to perform cell differentials on Wright-stained body fluids and to identify miscellaneous cells and inclusions in cytocentrifuged preparations using their own kit and result form. Participants will provide additional information about their competency assessment programs, continuing education, and professional background. Information will be collected from each site regarding their institution's minimum continuing education programs and requirements for their technologists in who review body fluids, and relevant procedures and policies related to body fluid review assessment.

Information will be collected from each site regarding their institution's minimum continuing education requirements for their technologists in hematology and relevant procedures and policies related to peripheral blood smear assessment.

Performance Indicators

- Individual technologist score (%) based on a standardized competency assessment method to determine a technologist's ability to identify various white blood cell types, microorganisms, and other cells and inclusions present in normal and abnormal cases in comparison to consensus responses
- Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your technical staff morphology and competency assessment requirements:

- · Result forms for up to 10 technologists (QPB10)
- Result forms for up to 25 technologists (QPB25)
- Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

*Participation in this study helps laboratories meet applicable requirements:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Checklist statements: HEM.35566, consistency of morphologic observation among personnel
 performing body fluid cell differentials at least annually; GEN.55500, Competency Assessment of Testing Personnel (element 5);
 GEN.55525, Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed
 by a qualified individual
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, Pl.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members

This is a one-time study conducted in the second quarter.

Assessment of Consistency of Peripheral Blood Morphologic Observations QPC10/QPC25

Introduction

The widespread use of automated white blood cell (WBC) differential counts and computer-generated whole slide imaging has decreased the time that the medical laboratory scientist/technologist staff dedicate to morphological assessment of blood cells. However, these staff must maintain their morphological skills. Laboratories have an annual requirement to do a morphologic comparison of their technical staff's peripheral blood smear results, assess their competency on peripheral blood smears, and provide appropriate education.

Objectives

This study will help assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the performance of accurate WBC differential counts and other peripheral blood smear morphological assessments. The evaluation provided will assist in the construction of individual educational programs for the technical staff and show areas that need focused review and improvement. The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for consistency of reporting morphology among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

A series of online, whole slide images of Wright or Wright-Giemsa stained peripheral blood smears using DigitalScope® technology will be available to each participating institution to assess technologists' performance on WBC differential counts and morphology assessment. Each technologist will receive their own kit. Technologists will provide information about their continuing education and professional background. Information will be collected from each site regarding their institution's minimum continuing education requirements for their technologists in hematology and relevant procedures and policies related to peripheral blood smear assessment.

Performance Indicators

- Individual technologist score (%) based on a standardized competency assessment method to determine a technologist's ability to identify various WBC types, red blood cell morphology, and platelet morphology in normal and abnormal cases
- · Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your staff technical competency assessment requirements:

- Result forms for up to 10 technologists (QPC10)
- Result forms for up to 25 technologists (QPC25)
- Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

Preliminary study reports are provided at institution and technologist levels.

*Participation in this study helps laboratories meet applicable requirements:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Checklist statements: HEM.34400, consistency of morphologic observation among personnel
 performing blood cell microscopy at least annually; GEN.55500, element 5, Competency Assessment of Testing Personnel; GEN.55525,
 Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed
 by a qualified individual
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, Pl.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members.

This is a one-time study conducted in the third quarter.

Assessment of Consistency of Gram Stain Morphologic Observations QPD10/QPD25

Introduction

Gram stain is a commonly performed bacterial stain in clinical microbiology laboratories. It is often the starting point guiding microbiological workup and initial clinical diagnosis and therapy. It is important for medical laboratory scientist/technologist staff who read Gram stains to provide an accurate interpretation based on reaction type and microscopic morphology, in order to provide presumptive identifications and quantification of bacteria and fungi in clinical specimens.

Objectives

This study will help assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the morphological assessment of Gram stains. Participation in this study will help management assess the technologist's ability to evaluate Gram stains using online, whole slide images. These cases provide a standardized review and evaluation for each technologist. The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for morphology consistency of reporting among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

A series of online, whole slide images of Gram-stained smears using DigitalScope technology will be provided to each participating institution to assess technologists' ability to detect various microorganisms. Technologists will provide information about their work experience related to Gram stains, continuing education, and professional background. Information will be collected from each laboratory site about their continuing education requirements in microbiology and relevant laboratory procedures and policies related to Gram stain assessment. Each technologist will receive their own kit and result form.

Performance Indicators

- Individual technologist score (%) for each Gram stain case, and overall based on a standardized competency assessment method
- · Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your staff technical competency assessment requirements:

- Result forms for up to 10 technologists (QPD10)
- Result forms for up to 25 technologists (QPD25)
- Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

Preliminary study reports are provided at institution and technologist levels.

*Participation in this study helps laboratories meet:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Microbiology Checklist statements: MIC.11060, Culture Result Reporting, personnel performing
 Gram stains for this purpose are subject to competency assessment; MIC.11350, Morphologic Observation Evaluation, the laboratory
 evaluates consistency of morphologic observation among personnel performing microscopic analysis (eg, stains, wet preparations) from
 direct specimens and cultured organisms at least annually. The laboratory director or designee must determine acceptability criteria
 for agreement.
- CAP Laboratory Accreditation Program Checklist items: GEN.55500, element 5, Competency Assessment of Testing Personnel; GEN.55525, Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed by a qualified individual.
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, Pl.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members

This is a one-time study conducted in the fourth quarter.

Continuous Quality Monitors

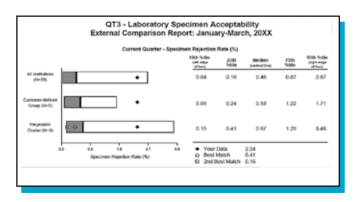
Use these programs to:

- · Identify and continuously monitor quality improvement over time.
- Measure the effectiveness and impact of implemented changes in key processes.

How It Works

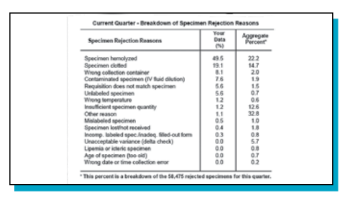
Step 1:

Establish realistic benchmarks by comparing your laboratory to others like yours.



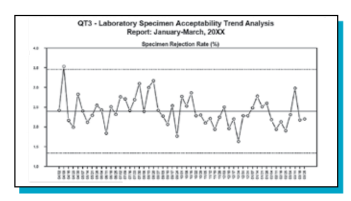
Step 2:

Identify improvement opportunities.



Step 3:

Monitor improvement over time to ensure accurate results, patient safety, and quality patient care.



The individual reports include performance of quality indicators over time, benchmarking information, trends, and suggested areas for improvement.

Participating laboratories receive:

- User Guide
- · Templates and instructions for data collection
- · Quarterly reports that include fingerprint clusters, customer-defined groups, and all-institution comparisons
- Access to the Peer Directory, allowing you to connect with your counterparts enrolled in the same program

Blood Culture Contamination QT2

Despite advances in blood culture practices and technology, false-positive blood culture results due to contaminants continue to be a critical problem. Blood culture contamination rate, the primary indicator of preanalytic performance in microbiology, is associated with increased length of hospital stay, additional expense, and the administration of unnecessary antibiotics.

The CAP and other accrediting organizations require you to monitor and evaluate key indicators of quality for improvement opportunities. Use this monitor to help meet CAP Laboratory Accreditation Program Checklist statements MIC.22630 and MIC.22635: "The laboratory must determine and regularly review the number of contaminated cultures. Tracking the contamination rate and providing feedback to units and persons drawing cultures is one method that has been shown to reduce contamination rates." This will also help laboratories meet The Joint Commission Standard QSA 04.07.01 EP3.

Objective

This study will determine the rate of blood culture contamination using standardized criteria for classifying contaminants.

Data Collection

On a monthly basis, participants will tabulate the total number of blood cultures processed and the total number of contaminated blood cultures. Blood cultures from neonatal patients are tabulated separately. For the purposes of this study, participants will consider a blood culture to be contaminated if they find one or more of the following organisms in only one of a series of blood culture specimens: Coagulase-negative *Staphylococcus*; *Micrococcus*; Alpha-hemolytic viridans group streptococci; *Propionibacterium acnes*; *Corynebacterium* sp. (diptheroids); or *Bacillus* sp. Participants have the option to monitor institution-specific subgroups (for example, a specific department or patient population).

Performance Indicators

- Neonatal contamination rate (%)
- Other contamination rate (%)
- Overall contamination rate (%)

Laboratory Specimen Acceptability QT3

A substantial amount of rework, diagnostic and therapeutic delay, and patient inconvenience can result from specimen rejection. Patient redraws may result from unlabeled, mislabeled, and incompletely-labeled specimens; clotted and/or hemolyzed specimens; or insufficient specimen quantity. By continuously monitoring specimen acceptability, collection, and transport, laboratories can promptly identify and correct problems. Enrollment in this study may assist the laboratory in monitoring compliance with CAP Laboratory Accreditation Program General Checklist statement GEN.40825: "There is a system to positively identify all patient specimens, specimen types, and aliquots at all times."

Objective

This study will identify and characterize unacceptable blood specimens that are submitted to the chemistry and hematology/coagulation sections of the clinical laboratory for testing.

Data Collection

This monitor includes all blood specimens submitted for testing to the chemistry and hematology departments of the clinical laboratory. On a weekly basis, participants will record the total number of specimens received, the number of rejected specimens, and the primary reason each specimen was rejected.

Performance Indicator

• Specimen rejection rate (%)

Performance Breakdown

• Breakdown of reasons for rejection (%)

In-Date Blood Product Wastage QT4

Blood for transfusion is a precious resource. At a minimum, wastage of blood that is not out-of-date represents a financial loss to the health care system. More ominously, systemic wastage of blood may reflect an environment of care that is out of control and may pose risks to patient safety.

Enrollment in this program assists laboratories in meeting regulatory requirements as follows:

- CAP Laboratory Accreditation Program Checklist statements TRM.40875, which requires the transfusion service medical
 director to monitor and audit transfusion practices to ensure the appropriate use of blood; TRM.30800, Disposition
 Records; and TRM.32275, Component Records, regarding recording the use of each blood or component product from
 receipt to final disposition.
- The Joint Commission Standards QSA.05.02.01, adequate blood and blood components; QSA.05.03.03, requirements for policies and procedures for returning unused blood products to blood transfusion services; and QSA.05.22.01, records of blood product disposition.
- AABB Standards for Blood Banks and Transfusion Services assessment 8.2, which requires transfusing facilities to have a peer-review program that monitors transfusion practices for blood components.

Objective

This study will compare the rates of blood product wastage (ie, units discarded in-date) in participating hospitals and track rates of improvement over time.

Data Collection

On a monthly basis, participants will use blood bank records to obtain information on the total number of units transfused for each type of blood component. Participants will track the number and type of blood units that are wasted in-date and the circumstances of wastage. This monitor includes the following types of blood components: whole blood (allogeneic), red blood cells (allogeneic), frozen plasma, platelet concentrates, single donor platelets, and cryoprecipitate.

Performance Indicators

• Overall blood wastage rate (%)

• Wastage rates by blood component type (%)

Performance Breakdown

• Breakdown of circumstances of wastage (%)

Satisfaction With Outpatient Specimen Collection QT7

Specimen collection is one of the few areas of laboratory medicine that involves direct outpatient contact. As a result, patient satisfaction with this service is a vital indicator of quality laboratory performance. The CAP's Laboratory Accreditation Program requires measurement of patient satisfaction with laboratory services (Checklist statement GEN.20335). Use this monitor to help meet this requirement.

Objective

This study will assess patient satisfaction with outpatient phlebotomy services by measuring patients' assessments of laboratory service hours, waiting time, comfort level, professionalism and courtesy, and privacy.

Data Collection

On a monthly basis, participants will provide copies of a standardized questionnaire in English and Spanish to a minimum of 25 outpatients (maximum of 99 outpatients) using predetermined data collection criteria. This monitor includes any outpatient undergoing venipuncture. It excludes patients seen in the emergency department, ambulatory surgery area, urgent care facility, chest pain center, 23-hour short-stay facility, employee health department, outpatient health screening fair/promotion, dialysis center, nursing home, or extended care facility.

Performance Indicators

- Satisfaction scores and satisfaction rates (% of patients rating 4 or 5) for the following categories:
 - o Overall experience
 - o Waiting time
 - o Patient comfort

- o Professionalism and courtesy
- o Patient privacy
- o Laboratory hours of operation

Stat Test Turnaround Time Outliers QT8

The stat test turnaround time (TAT) outlier rate, expressed as a percentage of tests missing target reporting times, is a measure of outcomes that evaluates how well the laboratory meets patient and clinician needs. This monitor helps meet CAP Laboratory Accreditation Program Checklist statement GEN.20316: "The QM program includes monitoring key indicators of quality in the preanalytic, analytic, and postanalytic phases."

Objective

This study will monitor the frequency that stat test TAT intervals exceed institutional stat test TAT expectations.

Data Collection

Before beginning data collection, participants will establish a specimen receipt-to-report deadline for emergency department (ED) stat potassium tests. On six predetermined days per month, participants will monitor the TAT of up to 10 randomly selected ED stat potassium tests on each of the three eight-hour shifts (up to 180 tests per month) and track the number of ED stat potassium results reported later than the established reporting deadline. This monitor includes stat potassium tests ordered as part of a panel and excludes stat potassium levels that are requested on body fluids other than blood, as part of timed or protocol studies, or after the specimen arrives in the laboratory.

Performance Indicator

Stat test TAT outlier rate (%)

Performance Breakdowns

- Breakdown of outliers by shift (%)
- Breakdown of outliers by day of week (%)

Critical Values Reporting QT10

Laboratories commonly refer to critical values as results requiring immediate notification to the physician or caregiver for necessary patient evaluation or treatment. Regulations from agencies and accreditors such as the CMS, The Joint Commission (National Patient Safety Goal NPSG.02.03.01), and the CAP Laboratory Accreditation Program (Checklist statement GEN.20316, COM.30000, COM.30100) mandate that laboratories develop and implement an alert system for critical values. Use this monitor to document compliance with your laboratory's alert plan.

Objective

This study will evaluate the documentation of successful critical values reporting in the general laboratory for inpatients and outpatients.

Data Collection

On a monthly basis, participants will evaluate 120 inpatient and 120 outpatient critical values. Data collection will include general chemistry, hematology, and coagulation analytes on the critical values list. Retrospectively, participants will record the total number of critical values monitored and the number with documentation of successful notification. In addition, participants will provide the number of critical values that were not communicated within three hours, the number of failed notifications due to laboratory oversight, and the number of successful notifications to licensed caregivers. This monitor will exclude critical values for cardiac markers, drugs of abuse, therapeutic drug levels, urinalysis, blood gases, point-of-care tests, and tests performed at reference laboratories.

Performance Indicators

- Total critical values reporting rate (%)
- Inpatient critical values reporting rate (%)
- Outpatient critical values reporting rate (%)
- Failed notification (< 3 hours) rate (%)

Corrected Results QT16

The CAP developed this monitor in recognition of the importance of timely detection and correction of erroneous laboratory results. Accuracy in laboratory results is critical to the effectiveness of a physician's plan of care for a patient. An erroneous result can delay or alter patient treatment; therefore, detection of erroneous results should be a priority in every laboratory and should be monitored as a key quality indicator. Use this monitor to help measure your compliance with CLIA 493.1299, Postanalytic Systems Quality Assessment, and help meet CAP Laboratory Accreditation Program Checklist statements GEN.20316, 41310, 41312, and The Joint Commission standard 02.12.01, Elements of Performance 9 and 10.

Objective

This study will monitor the number of corrected test results within individual institutions and compare performance with that of all institutions and those institutions similar to yours.

Data Collection

On a monthly basis, participants will monitor the number of corrected test results and the total number of billable tests for that month. Include test results for all patients in all care settings with the following exclusions: anatomic pathology tests, narrative physician-interpreted tests (eg, bone marrow biopsies and peripheral smear reports), and point-of-care tests.

Performance Indicator

• Test result correction rate (per 10,000 billable tests)

Outpatient Order Entry Errors QT17

Order accuracy bears an obvious relationship to the quality of laboratory testing. When the laboratory fails to complete a requested test, it delays the diagnostic evaluation, consumes unnecessary resources, causes patient inconvenience, and may prolong therapy. When the laboratory completes a test that was not requested, the cost of care increases, patients may be subjected to unnecessary phlebotomy, and laboratory efficiency declines. Use this monitor to help meet CAP Laboratory Accreditation Program Checklist statements GEN.20316, 40700, 40725, 40750 for test order and related information accuracy, and The Joint Commission Standard DC.01.02.01: The laboratory performs testing based on written laboratory test orders.

Objective

This study will measure the incidence of incorrectly interpreted and entered outpatient physician test orders into the laboratory information system, compare performance across institutions, and track performance over time.

Data Collection

On six preselected weekdays per month, participants will compare eight outpatient requisitions or order sheets to the orders entered into the laboratory's information system to determine if any order entry errors occurred.

This monitor includes test order review from ambulatory outpatients seen in offices and clinics operated by laboratory services, private physician offices, nursing homes, extended care facilities, and free-standing phlebotomy areas. Also included are send-out tests, chemistry, hematology, microbiology, immunology, toxicology, and urinalysis tests on outpatients. Order entry error categories include requesting physician errors, incorrect and extra test orders, missing test orders and diagnosis codes, test priority errors, and copy or fax result errors.

This monitor excludes tests performed in transfusion medicine or anatomic pathology, and also excludes tests from the following patient care settings: inpatient, emergency department, ambulatory surgery, urgent care, chest pain center, 23-hour short-stay facility, employee health department, outpatient screening fair/promotion, and dialysis center.

Performance Indicators

- Overall outpatient order entry error rate (%)
- Order entry error rates by type (%)

Performance Breakdown

• Breakdown of error types (%)

4

Quality Cross Check



Test multiple instruments at one time— Quality Cross Check is not PT and not subject to CMS restrictions.

Simplify biannual instrument comparability studies—receive customized reports that include peer group evaluations and instrument comparability statistics.

Discontinued Programs

Quality Cross Check — SARS-CoV-2, Serology (COVSQ)

Perform instrument comparability and stay in compliance

Quality Cross Check is a convenient solution to monitor instrument performance and assess comparability across multiple instruments in your laboratory and to identify potential issues before they affect patient results.

How It Works

- Receive three challenges in each of two mailings a year.
- Report up to three instruments for each challenge (and report up to 30 instruments for Quality Cross Check—Whole Blood Glucose).
- Receive a custom report package that includes peer group comparison and instrument comparability statistics for each reported analyte.

Stay in Compliance

In August 2015, the Centers for Medicare & Medicaid Services (CMS) reiterated that laboratories are not permitted to test proficiency testing (PT) samples on multiple instruments unless that is how the laboratory tests patient specimens.

The CMS interpretation was expanded beyond regulated analytes to include analytes not listed in Subpart I of the Clinical Laboratory Improvement Amendments regulations, including waived methods.

Quality Cross Check complements your existing CAP programs to monitor multiple instrument performance and is compliant with the CMS directive.

Monitoring Performance of Glucose Meters

Beginning in 2017, PT for waived whole blood glucose on glucose meters was no longer required for laboratories accredited by the CAP. Laboratories are required to perform alternative performance assessment.

In response to this change, the CAP introduced the Quality Cross Check—Whole Blood Glucose program (WBGQ). Participants in this program will enjoy the benefits of Quality Cross Check and have the ability to report up to 30 instruments for each challenge.

General Chemistry and Therapeutic Drug Monitoring

Quality Cross Check—Chemistry and Therapeutic Drug Monitoring CZQ					
Analyte Program Code Challenges per Shipment					
	CZQ				
See program CZ analytes on pages 54-56	ı	3			

This program does not meet regulatory requirements for proficiency testing; see program CZ on pages 54-56. For additional information about the Quality Cross Check program, see page 36.

Quality Cross Check—B-type Natriuretic Peptides BNPQ					
Analyte	Program Code Challenges per Shipment				
	BNPQ				
BNP	I	3			
NT-proBNP	I	3			

This program does not meet regulatory requirements for proficiency testing; see program BNP5 on page 59. For additional information about the Quality Cross Check program, see page 36.

Quality Cross Check—Whole Blood Glucose WBGQ						
Analyte	nalyte Program Code Challenges per Shipi					
	WBGQ					
Glucose		3				

The CAP Accreditation Programs require all accredited laboratories performing waived whole blood glucose testing using glucose meters to perform alternative performance assessment. This program can be used to meet alternative performance assessment requirements.

Program Information

- Three 5.0-mL liquid serum specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Program Information

- Three 1.5-mL liquid specimens
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

- Three 2.0-mL whole blood specimens
- Report up to 30 instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year



Quality Cross Check—Body Fluid Chemistry FLDQ				
Analyte	Program Code	Challenges per Shipment		
	FLDQ			
Albumin	I	3		
Amylase	I	3		
CA19-9	1	1		
Carcinoembryonic antigen (CEA)		1		
Cholesterol		3		
Creatinine		3		
Glucose		3		
Lactate		3		
Lactate dehydrogenase (LD)		3		
рН		3		
Protein, total		3		
Triglycerides		3		
Urea nitrogen	I	1		

This program does not meet regulatory requirements for proficiency testing; see program FLD on page 72. For additional information about the Quality Cross Check program, see page 36.

Quality Cross Check—Hemoglobin A1c GHQ					
Analyte	Program Code Challenges per Shipmen				
	GHQ				
Hemoglobin A1c	I	3			

This program does not meet regulatory requirements for proficiency testing; see program GH5 on page 62. For additional information about the Quality Cross Check program, see page 36.

Quality Cross Check—Cardiac Markers CRTQ					
Analyte Program Code Challenges per Shipme					
	CRTQ				
CK-MB, immunochemical		3			
Myoglobin		3			
Troponin I		3			

This program does not meet regulatory requirements for proficiency testing; see program CRT on page 60. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 3.0-mL simulated liquid body fluid specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Program Information

- Three 0.8-mL previously frozen liquid specimens in triplicate
- Report up to three instruments.
- Two shipments per year

- Three 2.0-mL liquid serum specimens
- Report up to three instruments.
- Two shipments per year

Quality Cross Check—High-Sensitivity Cardiac Markers HCRQ						
Analyte/Procedure Program Code Challenges per Shipmen						
HCRQ						
CK-MB, immunochemical	I	3				
Myoglobin	I	3				
High-sensitivity troponin I	I	3				
High-sensitivity troponin T ■ 3						

This program does not meet regulatory requirements for proficiency testing; see program HCRT on page 60. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 2.0-mL liquid specimens
- Report up to three instruments.
- Two shipments per year

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Julie Kingery, MD, FCAP Vice Chair of Clinical Pathology University of Florida

Endocrinology

Quality Cross Check-	Parathyroid Hor	rmone PTHQ
Analyte	Program Code	Challenges per Shipment
	PTHQ	
Parathyroid hormone (PTH)	I	3

This program does not meet regulatory requirements for proficiency testing; see program PTH on page 86. For additional information about the Quality Cross Check program, see page 36.

- Three 2.0-mL lyophilized serum specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Blood Gas, Critical Care, and Oximetry

Quality Cross Check—Blood Oximetry SOQ					
Analyte	Program Code Challenges per Shipme				
	SOQ				
Carboxyhemoglobin	I	3			
Hematocrit, estimated	I	3			
Hemoglobin, total	1	3			
Methemoglobin	I	3			
Oxyhemoglobin	I	3			

This program does not meet regulatory requirements for proficency testing; see program SO on page 95. For additional information about the Quality Cross Check program, see page 36.

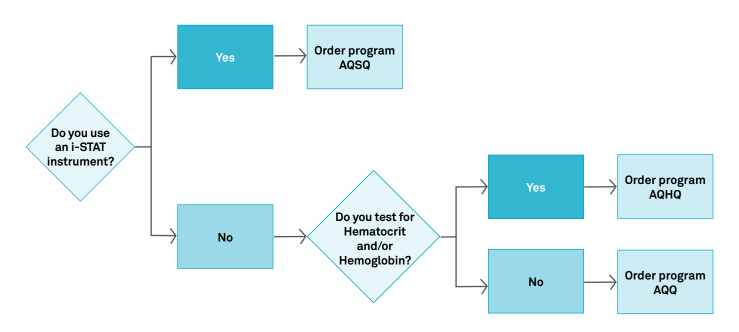
- Three 1.2-mL liquid specimens in triplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Quality Cross Check—Critical Care Blood Gas AQQ, AQHQ, AQSQ				
Analyte	F	Program Cod	Challenges per Shipment	
	AQQ	AQHQ	AQSQ	
Calcium, ionized	1			3
Chloride	1	1	•	3
Creatinine	1	•	•	3
Glucose	1		•	3
Hematocrit			•	3
Hemoglobin, estimated			•	3
Lactate	1			3
Magnesium, ionized	1	1		3
pCO ₂			•	3
pH	1			3
pO ₂			•	3
Potassium			•	3
Sodium	1	1	•	3
tCO ₂ (measured)				3
Urea nitrogen (BUN)	1	•	•	3

Additional Information

- It is not appropriate to report hemoglobin and hematocrit results by co-oximetry in these programs.
- These programs do no meet regulatory requirements for proficiency testing; see programs AQ, AQH, and AQIS on pages 92-93. For additional information about the Quality Cross Check program, see page 36.

- AQQ Three 2.5-mL specimens in triplicate; appropriate for all methods except i-STAT
- AQHQ Three 2.5-mL specimens in triplicate and three 2.5-mL specimens for hematocrit testing in triplicate; appropriate for all methods except i-STAT
- AQSQ Three 1.7-mL specimens in triplicate for i-STAT methods only
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year



Hematology and Clinical Microscopy

Quality Cross Check—Hematology FH3Q, FH4Q, FH9Q, FH13Q					
Analyte/Procedure	Program Code Challenges per Shipment				
	FH3Q	FH4Q	FH9Q	FH13Q	
Hematocrit	•	I		I	3
Hemoglobin	•	I		I	3
Immature granulocyte (IG)					3
Immature platelet fraction (IPF)%					3
Large unstained cells (LUC)	■ 3				
мсу, мсн, мснс	•	I		•	3
MPV	•	I		•	3
Nucleated red blood cell count (nRBC)					3
Platelet count	•	I	I	ı	3
RDW	I	I		I	3
Red blood cell count	I				3
WBC differential	I	ı		ı	3
White blood cell count	•			•	3

These programs do not meet regulatory requirements for proficiency testing; see the FH Series on page 140. For additional information about the Quality Cross Check program, see page 36.

- FH3Q, FH4Q, FH9Q, FH13Q -Three 2.5-mL whole blood specimens in vials with pierceable caps
- Report up to three instruments.
- For method compatibility, see instrument matrix on page 141.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Quality Cross Check—Reticulocyte RTQ, RT3Q, RT4Q Challenges per Instrument/Method **Program Code** Shipment RT4Q RTQ RT3Q Abbott Alinity hq, Abbott Cell-Dyn 4000, Sapphire, Siemens ADVIA 3 120/2120, and all other automated and manual methods Beckman Coulter, LH 500, LH 700 3 series, UniCel DxH series Sysmex XE-2100, XE-2100C, XE-2100D, XE-2100DC, XE-2100L, XE-5000, XN-L series, XN-series 3 (includes RL App), XR-series, XT-2000i, XT-4000i

These programs do not meet regulatory requirements for proficiency testing; see the RT Series on page 146. For additional information about the Quality Cross Check program, see page 36.

Quality Cross Check—Urinalysis CMQ			
Analyte	Program Code	Challenges per Shipment	
	СМQ		
Bilirubin	I	3	
Blood or hemoglobin	I	3	
Glucose		3	
hCG urine, qualitative	I	3	
Ketones	•	3	
Leukocyte esterase		3	
Nitrite		3	
Osmolality		3	
рН	I	3	
Protein, qualitative	I	3	
Reducing substances		3	

This program does not meet regulatory requirements for proficiency testing; see programs CMP and CMP1 on page 152. For additional information about the Quality Cross Check program, see page 36.

Program Information

- RTQ Three 1.0-mL stabilized red blood cell specimens
- RT3Q, RT4Q Three 3.0-mL stabilized red blood cell specimens
- Includes percentage and absolute result reporting
- Report up to three instruments.
- Two shipments per year

Program Information

- Three 10.0-mL liquid urine specimens for use with all instruments
- Report up to three instruments.
- Two shipments per year

Specific gravity

Urobilinogen

3

Quality Cross Check—Occult Blood OCBQ					
Analyte	Program Code Challenges per Shipr				
	OCBQ				
Occult blood		3			

This program does not meet regulatory requirements for proficiency testing; see program OCB on page 159. For additional information about the Quality Cross Check program, see page 36.

Program Information

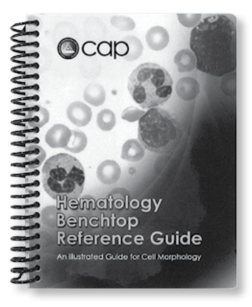
- Three 2.0-mL simulated fecal specimens
- Report up to three instruments.
- Two shipments per year

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Coagulation

Quality Cross Check-	—Coagulation CGL	Q
Analyte	Program Code	Challenges per Shipment
	CGLQ	
Activated partial thromboplastin time	I	3
Fibrinogen	I	3
Prothrombin time	I	3
D-dimer	I	2
Fibrin(ogen) degradation products, plasma	I	1
Fibrin(ogen) degradation products, serum	I	1

This program does not meet regulatory requirements for proficiency testing; see program CGL on page 166. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 1.0-mL lyophilized plasma specimens in triplicate, two 1.0-mL lyophilized plasma specimens, and one 2.0-mL serum specimen
- Report up to three instruments.
- · Two shipments per year

Quality Cross Check— Activated Clotting Time Series CTQ, CT1Q, CT2Q, CT3Q, CT5Q

Instrument/Cartridge		Program Code				Challenges per Shipment
	CTQ	CT1Q	CT2Q	CT3Q	CT5Q	
Helena Actalyke C-ACT®						3
Helena Actalyke MAX-ACT						
IL GEM Hemochron 100/ACT+						
IL GEM Hemochron 100/ACT-LR						
IL Hemochron® CA510/FTCA510						3
IL Hemochron FTK-ACT						3
IL Hemochron P214/P215						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT+						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT-LR						3
i-STAT Celite® and Kaolin ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus® HR-ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus LR-ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus R-ACT						3
Medtronic Hepcon HMS Plus		ı				3

These programs do not meet regulatory requirements for proficiency testing; see programs CT-CT3 and CT5 on page 170. For additional information about the Quality Cross Check program, see page 36.

- CTQ Three 3.0-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- CT1Q Three 1.7-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- CT2Q Three 0.5-mL lyophilized whole blood/ diluent ampules in triplicate
- CT3Q Three 0.5-mL lyophilized whole blood/ diluent ampules in triplicate
- CT5Q Three 1.7-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- Report up to three instruments.
- Two shipments per year

Microbiology

Quality Cross Check—SARS-CoV-2 Molecular COV2Q					
Analyte	Program Code Challenges per Ship				
	COV2Q				
SARS-CoV-2	I	3			

This program does not meet regulatory requirements for proficiency testing; see program COV2 on page 202. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 3.2-mL non-infectious liquid specimens that contain the whole SARS-CoV-2 genome
- Designed for molecular techniques
- Report up to three instruments.
- Two shipments per year

Quality Cross Check—SARS-CoV-2 Antigen COVAQ					
Analyte	Program Code	Challenges per Shipment			
	COVAQ				
SARS-CoV-2 antigen	I	3			

This program does not meet regulatory requirements for proficiency testing; see program COVAG on page 203. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 0.5-mL simulated respiratory specimens in triplicate
- Report up to three instruments.
- · Two shipments per year

Quality Cross Check—Nucleic Acid Amplification, Respiratory Limited ID3Q

,				
Analyte	Program Code	Challenges per Shipment		
	ID3Q			
Influenza A virus	ı	3		
Influenza B virus	ı	3		
Respiratory syncytial virus (RSV)	I	3		
SARS-CoV-2		3		

This program does not contain human genome material or sequences from human RNase P gene.

This program does not meet regulatory requirements for proficiency testing; see program ID3 on page 204. For additional information about the Quality Cross Check program, see page 36.

- Three 1.0-mL liquid specimens
- Designed for molecular multiplex panel users
- Report up to three instruments.
- · Two shipments per year

Transfusion Medicine

Quality Cross Check—Transfusion Medicine JATQ					
Procedure	Program Code	Challenges per Shipment			
	JATQ				
ABO grouping	•	3			
Antibody detection	I	3			
Rh typing	I	3			

This program does not meet regulatory requirements for proficiency testing; see program JAT on page 233. For additional information about the Quality Cross Check program, see page 36.

Program Information

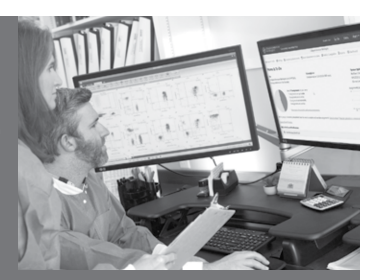
- Three 6.0-mL 13%-17% whole blood specimens
- May be used with automated and manual procedures
- · Two shipments per year

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5

Point-of-Care Programs



Keep your point-of-care (POC) instruments and staff operating at peak performance.

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- Gain insights with the Point-of-Care Testing Toolkit, an ebook resource for all members of the team.

Point-of-Care Programs

POC Competency Challenges help POC coordinators streamline operator education (initial training and ongoing competency). These programs include standardized specimens that can be used not only to train operators and assess competency, but also to evaluate/troubleshoot instrument and method performance for waived and non-waived tests.

Expected results will be provided. These programs are not proficiency testing programs and participants will not return results to the CAP.

POC Competency Challenges have limited availability and stability. These programs must be purchased by May 1.

POC Competency Challenges POC1, POC2, POC3, POC4					
Program Name	Program Code				Challenges per Shipment
	POC1	POC2	POC3	POC4	
hCG Competency					10
Glucose Competency					10
Urine Dipstick Competency					10
Strep Screen Competency				I	10

Program Information

- POC1 One positive 10.0-mL liquid urine specimen
- POC2 One abnormal 2.0-mL whole blood specimen
- POC3 One abnormal 10.0-mL liquid urine specimen
- POC4 One 1.0-mL positive liquid specimen
- Each program provides material to test up to 10 staff.

POC Competency Challenges POC6, POC7, POC8, POC9					
Program Name		Progra	Challenges per Shipment		
	POC6	POC7	POC8	POC9	
PT/INR, Roche CoaguChek Pro II, XS Plus, and XS Pro Competency					10
Waived Chemistry, Glucose, and Hemoglobin Competency					10
Influenza A/B Antigen Detection Competency					10
Fecal Occult Blood Competency				I	10

- POC6 One abnormal 0.3-mL lyophilized plasma specimen (five vials) and five corresponding diluents
- POC7 One abnormal 1.5-mL whole blood specimen compatible with the HemoCue® B, HemoCue 201, and Stanbio HemoPoint® H2 instruments
- POC8 One 1.5-mL positive liquid specimen for influenza A; one 1.5-mL positive liquid specimen for influenza B
- POC9 One positive 2.0-mL fecal specimen
- Each program provides material to test up to 10 staff.

POC Competency Challenges POC10, POC11, POC12					
Program Name		Program Code			
	POC10	POC11	POC12		
Blood Gases Competency	ı			10	
Blood Gases, i-STAT Competency				10	
Point-of-Care Cardiac Markers Competency			I	10	

- POC10 One abnormal 2.5-mL aqueous blood gas specimen (10 vials) and one 2.5-mL hematocrit/ hemoglobin specimen (10 vials)
- POC11 One abnormal 2.5-mL aqueous specimen (10 vials) for blood gas and hematocrit/hemoglobin testing
- POC12 One 1.5-mL plasma specimen (two vials); compatible with plasma-based tests, such as Alere Triage® and i-STAT instruments
- Each program provides material to test up to 10 staff.

Professionalism in Pathology and Laboratory Medicine

This important resource provides a basic understanding of how ethics and professionalism impact pathology and laboratory medicine. Approaches and guidance to educational and assessment tools, including more than 100 case vignettes to guide discussion, are included. The book also discusses professionalism in the context of research, pathologist wellbeing and burnout, legal aspects, diversity, organizational leadership, and patient safety and quality of care. Also addressed are lapses in ethical and professional behavior as well as recommendations on future directions for research and education in professionalism.

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POC Competency Challenges POC14, POC15, POC16					
Program Name		Program Code			
	P0C14	POC15	POC16		
Medtronic ACT/ACT Plus®, i-STAT Competency				5	
Hemochron® Jr., IL GEM PCL ACT-LR Competency		•		5	
Hemochron Jr., Signature, IL GEM PCL ACT Competency			1	5	

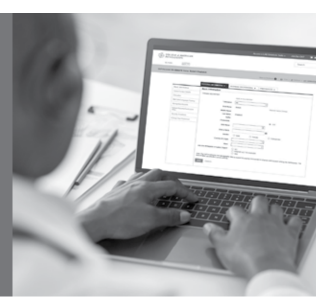
- POC14 Five abnormal
 1.7-mL lyophilized whole
 blood specimens with five
 corresponding diluents
 and one calcium chloride
 diluent vial; compatible with
 Medtronic Hemotec ACT/
 ACTII/ACT Plus, Medtronic
 Hepcon HMS/HMS Plus, and
 i-STAT Celine and Kaolin ACT
- POC15 Five abnormal 0.5-mL lyophilized whole blood/diluent ampules; compatible with IL GEM PCL Plus ACT-LR and ITC Hemochron Jr., Signature ACT-LR
- POC16 Five abnormal 0.5-mL lyophilized whole blood/diluent ampules; compatible with IL GEM PCL Plus ACT and ITC Hemochron Jr., Signature ACT+
- Each program provides material to test up to five staff.

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General Chemistry and Therapeutic Drug Monitoring



CAP Accreditation: Focused on the laboratory

CAP laboratory accreditation gives you and your staff the confidence of knowing that your laboratory is providing the highest quality results and better patient outcomes.

General Chemistry and Therapeutic Drug Monitoring

All current and new 2025 Centers for Medicare & Medicaid Services (CMS) regulated analytes are listed in **bold** type.

General Chemistry and Therapeutic Drug Monitoring	54
Urine Chemistry	
Special Chemistry	71
•	

New Programs



Point-of-Care High-Sensitivity Troponin I (PCHT)	65
Waived Hemoglobin (HCC1)	66

Program Changes

CAP/AACC immunosuppressive Drugs (CS) is now called	
CAP/ADLM Immunosuppressive Drugs (CS)	58
Hemoglobin A1c, 3 Challenge (GH2) is now called Hemoglobin A1c, Waived and is	
for waived methods only	62
High-Sensitivity C-reactive Protein (HSCRP) number of challenges, number of shipments	63

Discontinued Programs

B-type Natriuretic Peptides, 2 Challenge (BNP) Sweat Analysis (SW1)

General Chemistry and Therapeutic Drug Monitoring

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

General Chemistry and Therapeutic Drugs C1, C3/C3X, C4, CZ/CZX/CZ2X, Z

01,007000,04,0270227,2						
Analyte		Program Code			Challenges per Shipment	
	C1	C3/C3X	C4	CZ/CZX/ CZ2X	z	
Alanine aminotransferase (ALT/SGPT)	•	•		•		5
Albumin	ı					5
Alkaline phosphatase	ı					5
Amylase	ı					5
Aspartate aminotransferase (AST/SGOT)	ı			•		5
Bilirubin, direct	ı		ı			5
Bilirubin, total*	ı		ı	I		5
Calcium	•		ı			5
Chloride	ı		ı	I		5
Cholesterol, total	ı		ı	I		5
Cortisol	ı			I		5
Creatine kinase (CK)	1			I		5
Creatinine	ı		ı	I		5
Glucose	ı		ı			5
HDL cholesterol	ı		ı	I		5
Human chorionic gonadotropin (hCG), quantitative	ı			•		5
Iron	ı					5
Lactate dehydrogenase (LD)	ı			I		5
LDL cholesterol, measured	ı		ı	I		5
Lipoprotein (a)	•					5
Magnesium	ı			I		5
Pancreatic amylase	ı			I		5
Potassium	ı		ı			5
Protein, total	ı			I		5
Sodium	ı	•	ı	I		5
Triiodothyronine (T3), free	•	•		I		5
Triiodothyronine (T3), total	•	•		•		5
T3, uptake and related tests	ı					5

Continued on the next page

- C1, C3, C4, CZ, Z Five 5.0-mL liquid serum specimens
- C3X, CZX Five 5.0-mL liquid serum specimens in duplicate
- CZ2X Five 5.0-mL liquid serum specimens in triplicate
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year
- For multiple instrument reporting options, see the Quality Cross Check program, CZQ, on page 56.



^{*}General Chemistry and Therapeutic Drugs programs do not fulfill the neonatal bilirubin proficiency testing requirements for the CAP Laboratory Accreditation Programs. See programs NB, NB2, on page 64.

General Chemistry and Therapeutic Drugs C1, C3/C3X, C4, CZ/CZX/CZ2X, Z continued

C1, C3/C3X, C4,	CZ/	CZX/C	ZZ)	⟨,∠ contir	nued	
Analyte					Challenges per Shipment	
	C1	C3/C3X	C4	CZ/CZX/ CZ2X	z	
Thyroxine (T4), free	I			ı		5
Thyroxine (T4), total						5
Thyroid-stimulating hormone (TSH)	•			•		5
Triglycerides				ı		5
Urea nitrogen (BUN)				ı		5
Uric acid				ı		5
Acid phosphatase				ı		5
Ammonia				ı		5
Apolipoprotein A1				I		5
Apolipoprotein B				I		5
Calcium, ionized				ı		5
Carbon dioxide (CO ₂)				I		5
Ferritin				I		5
Gamma glutamyl transferase (GGT)				ı		5
Iron binding capacity, total (measured)				ı		5
Iron binding capacity, unsaturated (measured)				ı		5
Lactate				ı		5
Lipase				ı		5
Osmolality				ı		5
Phosphorus				ı		5
Prealbumin				I		5
Transferrin				ı		5
Lithium				ı	ı	5
Acetaminophen				I	ı	5
Amikacin				ı	ı	5
Caffeine				ı	ı	5
Carbamazepine				I	ı	5
Carbamazepine, free				ı	1	5
Digoxin				ı	ı	5
Digoxin, free				ı	•	5
Disopyramide				ı	ı	5
Continued on the next page						

- C1, C3, C4, CZ, Z Five 5.0-mL liquid serum specimens
- C3X, CZX Five 5.0-mL liquid serum specimens in duplicate
- CZ2X Five 5.0-mL liquid serum specimens in triplicate
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year
- For multiple instrument reporting options, see the Quality Cross Check program, CZQ, on page 56.



Vancomycin

General Chemistry and Therapeutic Drugs C1, C3/C3X, C4, CZ/CZX/CZ2X, Z continued Challenges per **Program Code Analyte** Shipment CZ/CZX/ C1 C3/C3X Z C4 CZ2X Ethosuximide ı 5 Gentamicin 5 Lidocaine 5 Methotrexate 5 N-acetylprocainamide (NAPA) 5 Phenobarbital 5 Phenytoin 5 Phenytoin, free 5 Primidone 5 Procainamide 5 Quinidine 5 5 Salicylate 5 Theophylline 5 Tobramycin 5 Valproic acid 5 Valproic acid, free ı

Program Information

- C1, C3, C4, CZ, Z Five 5.0-mL liquid serum specimens
- C3X, CZX Five 5.0-mL liquid serum specimens in duplicate
- CZ2X Five 5.0-mL liquid serum specimens in triplicate
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year
- For multiple instrument reporting options, see the Quality Cross Check program, below.



5

Quality Cross Check—Chemistry and Therapeutic Drug Monitoring CZQ Analyte Program Code Challenges per Shipment CZQ See program CZ analytes on pages 54-56

This program does not meet regulatory requirements for proficiency testing; see program CZ on pages 54-56. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

- Three 5.0-mL liquid serum specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Accuracy-Based Lipids ABL				
Analyte	Program Code	Challenges per Shipment		
	ABL			
Apolipoprotein A1	I	3		
Apolipoprotein B	I	3		
Cholesterol*	I	3		
HDL cholesterol*	I	3		
Non-HDL cholesterol	I	3		
LDL cholesterol	ı	3		
Lipoprotein(a)	I	3		
Triglycerides*	1	3		

^{*}This analyte will be evaluated against the reference method.

Harmonized Thyroid ABTH				
Analyte	Program Code	Challenges per Shipment		
	ABTH			
Triiodothyronine (T3), free	1	3		
Triiodothyronine (T3), total	ı	3		
Thyroxine (T4), free	ı	3		
Thyroxine (T4), total	1	3		
Thyroid-stimulating hormone (TSH)	ı	3		

Additional Information

- Analytes will be evaluated using harmonization.
- Specimens are collected by a modified application of Clinical and Laboratory Standards Institute Guideline CLSI C37-A, Preparation and Validation of Commutable Frozen Human Serum Pools as Secondary Reference Materials for Cholesterol Measurement Procedures; Approved Guideline.

Thyroid Calibration Verification	.N50	
Analyte	Program Code	LN50 Target Ranges
	LN50	
Triiodothyronine (T3), free	•	1.0-18.0 pg/mL
Triiodothyronine (T3), total	ı	0.4-7.0 ng/mL
Thyroxine (T4), free	I	0.7–7.0 ng/dL
Thyroxine (T4), total	ı	1.0-27.0 μg/dL
Thryoid-stimulating hormone (TSH)	I	0.1–120.0 μIU/mL

Program Information

- Three 1.0-mL human serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Program Information

- Three 1.0-mL frozen human serum specimens
- Two shipments per year

- Eighteen 2.0-mL serum specimens
- · Two shipments per year

CAP/ADLM Immunosuppressive Drugs CS				
Analyte	Program Code	Challenges per Shipment		
	CS			
Cyclosporine		3		
Sirolimus (rapamycin)		3		
Tacrolimus		3		

- Three 5.0-mL whole blood specimens
- For laboratories monitoring cyclosporine, sirolimus, and tacrolimus in transplant patients
- Two shipments per year



Antifungal Drugs Monitoring AFD				
Procedure	Program Code	Challenges per Shipment		
	AFD			
Fluconazole	I	3		
Itraconazole	I	3		
Posaconazole	I	3		
Voriconazole	I	3		

Program Information

- Three 2.0-mL serum specimens
- For laboratories performing quantitative analysis of antifungal agents
- Two shipments per year

Everolimus EV				
Analyte	Program Code	Challenges per Shipment		
	EV			
Everolimus	1	3		

Program Information

- Three 4.0-mL whole blood specimens
- Two shipments per year

Mycophenolic Acid MPA					
Analyte	Program Code	Challenges per Shipment			
	MPA				
Mycophenolic acid		3			

- Three 5.0-mL lyophilized serum specimens
- Two shipments per year

Therapeutic Drug Monitoring—Extended ZE				
Analyte	Program Code	Challenges per Shipment		
	ZE			
Clozapine	•	3		
Gabapentin	•	3		
Lacosamide	•	3		
Lamotrigine	•	3		
Levetiracetam	•	3		
Oxcarbazepine	I	3		
Oxcarbazepine metabolite		3		
Pregabalin	•	3		
Rufinamide	I	3		
Teriflunomide	•	3		
Topiramate	•	3		
Zonisamide		3		

- Three 5.0-mL serum specimens
- Two shipments per year

Therapeutic Drug Monitoring—Special ZT			
Analyte	Program Code	Challenges per Shipment	
	ZT		
Amitriptyline	•	3	
Desipramine	ı	3	
Imipramine	I	3	
Nortriptyline	1	3	
Tricyclics, total (qualitative/ quantitative)		3	

Program Information

- Three 5.0-mL lyophilized serum specimens
- Two shipments per year

B-type Natriuretic Peptides BNP5			
Analyte	Program Code	Challenges per Shipment	
	BNP5		
BNP	I	5	
NT-proBNP	I	5	

Additional Information

- For i-STAT, Quidel Triage, and Pathfast, use Point-of-Care Cardiac Markers programs PCARM or PCARMX.
- For second instrument reporting options, see the Quality Cross Check program, BNPQ, on page 60.

- Five 1.0-mL liquid plasma specimens
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year

Quality Cross Check—B-type Natriuretic Peptides BNPQ			
Analyte	Program Code Challenges per Shipment		
	BNPQ		
BNP	I	3	
NT-proBNP	I	3	

This program does not meet regulatory requirements for proficiency testing; see program BNP5, on page 59. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Cardiac Markers CRT, CRTI, HCRT, HCRTI					
Analyte	Program Code			Challenges per Shipment	
	CRT	CRTI	HCRT	HCRTI	
CK-MB, immunochemical					5
CK isoenzymes (CK-BB, CK-MB , CK-MM), electrophoretic				•	5
LD1, LD2, LD3, LD4, LD5, electrophoretic					5
LD1/LD2 ratio calculation and interpretation					5
Myoglobin	•				2
Troponin I					5
High-sensitivity troponin I					5
High-sensitivity troponin T					5

Program Information

- Three 1.5-mL liquid specimens
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

- CRT Five 2.0-mL liquid specimens
- CRTI Ten 2.0-mL liquid specimens
- HCRT Five 2.0-mL liquid specimens
- HCRTI Ten 2.0-mL liquid specimens
- Three shipments per year

Quality Cross Check—High-Sensitivity Cardiac Markers HCRQ				
Analyte/Procedure Program Code Challenges per Shipme				
HCRQ				
CK-MB, immunochemical	I	3		
Myoglobin	I	3		
High-sensitivity troponin I	I	3		
High-sensitivity troponin T	I	3		

This program does not meet regulatory requirements for proficiency testing; see program HCRT, on page 60. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Quality Cross Check—Cardiac Markers CRTQ			
Analyte	Program Code	Challenges per Shipment	
	CRTQ		
CK-MB, immunochemical		3	
Myoglobin	I	3	
Troponin I	I	3	

This program does not meet regulatory requirements for proficiency testing; see program CRT, on page 60. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Program Information

- Three 2.0-mL liquid specimens
- Report up to three instruments.
- · Two shipments per year

- Three 2.0-mL liquid serum specimens
- Report up to three instruments.
- · Two shipments per year

Hemoglobin A1c Waived GH2					
Analyte	Analyte Program Code Challenges per Shipmen				
GH2					
Hemoglobin A1c ■ 3					

Additional Information

- This program will be evaluated against the National Glycohemoglobin Standardization Program (NGSP) reference method.
- For multiple instrument reporting options, see the Quality Cross Check program, GHQ, below.
- This program has limited stability. Laboratories outside the US or Canada should consider purchasing GH5I, which has longer stability.

Program Information

- · Accuracy-Based program
- Three 0.8-mL liquid human whole blood specimens
- Two shipments per year
- Designed for waived methods

Hemoglobin A1c GH5			
	Program Code	Challenges per Shipment	
	GH5		
Hemoglobin A1c		5	

Additional Information

- This program will be evaluated against the National Glycohemoglobin Standardization Program (NGSP) reference method.
- For multiple instrument reporting options, see the Quality Cross Check program, GHQ, below.
- This program has limited stability. Laboratories outside the US or Canada should consider purchase of GH5I, which has longer stability.

Quality Cross Check—Hemoglobin A1c GHQ			
Analyte	Program Code Challenges per Shipment		
	GHQ		
Hemoglobin A1c	I	3	

This program does not meet regulatory requirements for proficiency testing; see program GH5, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Program Information

- Accuracy-Based program
- Five 0.8-mL liquid human whole blood specimens
- · Three shipments per year

- Three 0.8-mL previously frozen liquid specimens in triplicate
- Report up to three instruments.
- · Two shipments per year

Hemoglobin A1c GH5I			
Analyte	Program Code Challenges per Shipmen		
	GH5I		
Hemoglobin A1c		5	

This program will not be evaluated against the National Glycohemoglobin Standardization Program (NGSP) reference method. See program GH5 to be evaluated against the NGSP reference method.

Program Information

- Five 0.5-mL lyophilized specimens with a 3.0-mL dropper-tipped vial of diluent
- Designed for laboratories outside the US that have experienced significant shipping and receiving issues and require longer specimen stability
- · Three shipments per year

Glycated Serum Albumin GSA					
Analyte Program Code Challenges per Shipr					
GSA					
Glycated serum albumin	•	3			

Program Information

- Three 1.0-mL liquid serum specimens
- Two shipments per year

High-Sensitivity C-reactive Protein HSCRP			
Analyte Program Code Challenges per Shipmen			
	HSCRP		
High-sensitivity C-reactive protein	ı	5	

Program Information

- Five 0.5-mL liquid serum specimens
- Three shipments per year

Homocysteine HMS			
Analyte Program Code Challenges per Shipm			
	HMS		
Homocysteine	I	3	

Program Information

- Three 1.0-mL serum specimens
- Two shipments per year

Ketones KET			
Analyte	Program Code	Challenges per Shipment	
	KET		
Beta-hydroxybutyrate	I	2	
Total ketones	I	2	

- Two 2.0-mL serum specimens
- For semi-quantitative methods using the nitroprusside reaction for total ketones testing
- Two shipments per year

Chemistry—Limited, Waived LCW			
Analyte	Program Code	Challenges per Shipment	
	LCW		
Cholesterol		3	
Glucose		3	
HDL cholesterol		3	
LDL cholesterol		3	
Triglycerides		3	

- Three 3.0-mL liquid serum specimens
- For use with waived methods such as the Cholestech LDX® and Roche Accu-Chek® Instant Plus
- The glucose specimens are not appropriate for use on other whole blood glucose meters.
- Two shipments per year

Neonatal Bilirubin NB, NB2			
Analyte Challenges per Shipment			
	Program Code		
	NB NB2		
Bilirubin, direct	2	2	
Bilirubin, total 5 2			

One human-based serum specimen will offer the value assigned using the reference method procedure (*Clin Chem.* 1985;31:1779-1789).

Program Information

- NB Five 1.0-mL human serum specimens; three shipments per year
- NB2 Two 1.0-mL human serum specimens; must order in conjunction with a five-challenge total bilirubin general chemistry proficiency testing program to meet regulatory requirements; two shipments per year
- Conventional and International System of Units (SI) reporting offered

Point-of-Care Cardiac Markers PCARM/PCARMX			
Analyte	Program Code Challenges per Shipment		
	PCARM PCARMX		
BNP	ı		5
СК-МВ	ı		5
D-dimer	ı		2
Myoglobin	ı		2
NT-proBNP			5
Troponin I	ı		5

- PCARM Five 1.5-mL liquid EDTA plasma specimens for point-of-care instruments such as Quidel Triage, Pathfast, and i-STAT
- PCARMX All PCARM specimens in duplicate
- Three shipments per year

Point-of-Care High-Sensitivity Troponin I PCHT			
Analyte Program Code Challenges per Shipment			
PCHT			
High-sensitivity troponin I	I	5	

- Five 1.5-mL liquid specimens for point-of-care instruments such as i-STAT
- Three shipments per year

Whole Blood Chemistry Compatibility Matrix

Whole Blood Analyzer/Method	Analyte	Compatible Survey Programs	Page
HemoCue® Glucose 201 systems	Glucose	нсс	below
HemoCue Hb 201+ systems	Hemoglobin	HCC	below
HemoCue Hb 301 and 801 systems	Hemoglobin	HCC1	below
Roche Reflotron®	Cholesterol	C1 C4	54-56
	Glucose	C1, C4	54-56
Cholestech LDX®	Total cholesterol		64
	HDL cholesterol	1.004	64
	Triglycerides	LCW	64
	Glucose		64
Whole blood cholesterol meters	Cholesterol	C1, C4, LCW	54, 64
Whole blood glucose meters	Glucose	HCC2, WBGQ	66.67
Nova StatSensor®/ StatSensor Xpress™	Creatinine	WBCR	66

Waived Combination HCC			
Analyte Program Code Challenges per Shipment			
	HCC		
Hemoglobin		2	
Whole blood glucose		2	

- Two 1.5-mL whole blood specimens; two shipments per year
- Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix above.

Waived Hemoglobin HCC1			
Analyte	Program Code	Challenges per Shipment	
	HCC1		
Hemoglobin		2	

- Two 1.0-mL whole blood specimens; two shipments per year
- Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix on page 65.

Waived Combination HCC2			
Analyte	Program Code	Challenges per Shipment	
	HCC2		
Hematocrit		2	
Hemoglobin	I	2	
Urinalysis/urine hCG		2	
Whole blood glucose		3	

Program Information

- Total of four shipments per year
- Hematocrit, hemoglobin, and urinalysis/urine hCG testing
 Two 3.0-mL whole blood specimens and two 10.0-mL urine specimens; two shipments per year: A and C
- Whole blood glucose testing

 Three 2.0-mL whole blood specimens; two shipments per year: B and D
- Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix on page 65.

Whole Blood Creatinine WBCR				
Analyte	lyte Program Code Challenges per Shipme			
	WBCR			
Creatinine		5		

- Five 4.0-mL whole blood specimens
- For use with the Nova StatSensor/StatSensor Xpress
- Three shipments per year

Quality Cross Check—Whole Blood Glucose WBGQ			
Analyte	Program Code	Challenges per Shipment	
	WBGQ		
Glucose		3	

The CAP Accreditation Programs require all accredited laboratories performing waived whole blood glucose testing using glucose meters to perform alternative performance assessment. This program can be used to meet alternative performance assessment requirements.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- · Identify and troubleshoot instrument/method problems.
- · Correlate results with other laboratories or instruments.
- Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- · Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Chemistry/TDM, Validated Material

Validated Material	Program Code	Corresponding Program	Pages
Chemistry/TDM	CZVM	CZ	54-56

Program Information

- Three 2.0-mL whole blood specimens
- Report up to 30 instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year



Program Information

• Five 5.0-mL liquid serum specimens

Urine Chemistry

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Urine Chemistry—General U			
Analyte	Program Code	Challenges per Shipment	
	U		
Amylase	I	3	
Calcium	I	3	
Chloride	I	3	
Creatinine	I	3	
Glucose	I	3	
Magnesium	1	3	
Nitrogen, total	I	3	
Osmolality	I	3	
Phosphorus	I	3	
Potassium	I	3	
Protein, total	I	3	
Sodium	1	3	
Urea nitrogen	I	3	
Uric acid	I	3	
Urine albumin, quantitative	I	3	
Urine albumin:creatinine ratio	ı	3	

Program Information

- Six 15.0-mL urine specimens
- One mailing per year will include an additional educational specimen for uric acid testing for a total of seven challenges per year.
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Accuracy-Based Urine ABU			
Analyte	Program Code	Challenges per Shipment	
	ABU		
Calcium		3	
Creatinine	I	3	
Protein, total		3	
Urine albumin, quantitative	ı	3	
Urine albumin: creatinine ratio	I	3	

- Three 5.0-mL human urine specimens
- Two shipments per year

Kidney Stone Risk Assessment KSA			
Analyte	Program Code	Challenges per Shipment	
	KSA		
Citrate	ı	3	
Cystine	ı	3	
Oxalate		3	

- Three 13.5-mL liquid urine specimens
- Two shipments per year

Urine Chemistry—Special N			
Analyte	Program Code	Challenges per Shipment	
	N		
3-methoxytyramines		3	
5-hydroxyindoleacetic acid	I	3	
17-hydroxycorticosteroids		3	
17-ketosteroids		3	
Aldosterone	I	3	
Coproporphyrins	I	3	
Cortisol, urinary free	I	3	
Dopamine	I	3	
Epinephrine	I	3	
Homovanillic acid	I	3	
Metanephrine	I	3	
Norepinephrine		3	
Normetanephrine	I	3	
Uroporphyrin	I	3	
Vanillylmandelic acid	I	3	

Program Information

- N Six 10.0-mL lyophilized urine specimens and three 10.0-mL liquid urine specimens
- Two shipments per year

Myoglobin, Urine MYG			
Analyte	Program Code	Challenges per Shipment	
	MYG		
Myoglobin, urine, qualitative and quantitative	ı	2	

- Two 1.0-mL urine specimens
- Two shipments per year

Porphobilinogen, Urine UPBG			
Analyte	Program Code	Challenges per Shipment	
	UPBG		
Porphobilinogen		3	

- Three 5.0-mL urine specimens
- For use with qualitative and quantitative methods
- · Two shipments per year

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- Identify and troubleshoot instrument/method problems.
- · Correlate results with other laboratories or instruments.
- Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Urine Chemistry—General, Validated Material

Validated Material	Program Code	Corresponding Program	Page
Urine Chemistry	UVM	U	68

Program Information

- Six 15.0-mL urine specimens
- One mailing per year will include an additional specimen for uric acid testing.

The CAP is your trusted calibration verification and linearity partner, offering a comprehensive menu of programs for diagnostic confidence.

- **Expedited results**—View your linearity evaluation for most CVL programs within two business days of data submission.
- Customized report package—Let our team of biostatisticians perform the statistical analysis of your results so you don't have to.
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See the Instrumentation Verification Tools section of this catalog to determine programs that best fit your laboratory's CVL needs.

Special Chemistry

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

1,5-Anhydroglucitol AG			
Analyte Program Code Challenges per Shipm			
	AG		
1,5-anhydroglucitol	I	3	

Program Information

- Three 1.0-mL liquid serum specimens
- Two shipments per year

Aldolase ADL			
Analyte Program Code Challenges per Shipme			
	ADL		
Aldolase	I	2	

Program Information

- Two 3.0-mL liquid serum specimens
- Two shipments per year

Angiotensin Converting Enzyme ACE			
Analyte Program Code Challenges per Shipn			
	ACE		
Angiotensin converting enzyme, quantitative		2	

- Two 2.0-mL lyophilized serum specimens
- Two shipments per year

Body Fluid Chemistry FLD			
Analyte	Program Code	Challenges per Shipment	
	FLD		
Albumin	1	3	
Amylase	1	3	
CA19-9	1	1	
CEA	I	1	
Cholesterol	1	3	
Creatinine	1	3	
Glucose	1	3	
Lactate	1	3	
Lactate dehydrogenase (LD)	I	3	
рН	I	3	
Protein, total	I	3	
Triglycerides	1	3	
Urea nitrogen	1	1	

- Three 3.0-mL simulated liquid body fluid specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

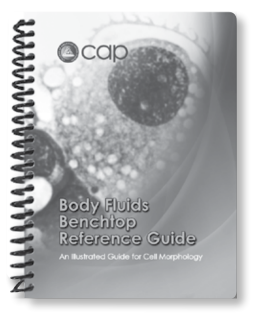
For multiple instrument reporting options, see the Quality Cross Check program, FLDQ, on page 73.

Body Fluids Benchtop Reference Guide

- Thirty-six color images, including common and rare cells, crystals, and other cell inclusions
- Detailed descriptions of each cell including facts, cell morphology, and inclusions
- Nine tabbed sections for easy reference
 - Erythroid Series
 - o Lymphoid Series
 - Myeloid Series
 - Mononuclear Phagocytic Series
 - Lining Cells
 - Miscellaneous Cells
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Quality Cross Check—Body Fluid Chemistry FLDQ			
Analyte	Program Code	Challenges per Shipment	
	FLDQ		
Albumin	I	3	
Amylase	I	3	
CA19-9	I	1	
Carcinoembryonic antigen (CEA)	I	1	
Cholesterol	I	3	
Creatinine	I	3	
Glucose	I	3	
Lactate	I	3	
Lactate dehydrogenase (LD)	I	3	
рН	I	3	
Protein, total		3	
Triglycerides	I	3	
Urea nitrogen	I	1	

This program does not meet regulatory requirements for proficiency testing; see program FLD, on page 72. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Body Fluid Chemistry 2 FLD2			
Analyte	Program Code	Challenges per Shipment	
	FLD2		
Alkaline phosphatase		3	
Bilirubin	1	3	
Calcium	I	3	
Chloride	I	3	
Lipase	I	3	
Potassium	I	3	
Sodium	I	3	
Uric acid	ı	3	

Program Information

- Three 3.0-mL simulated liquid body fluid specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

- Three 3.0-mL liquid body fluid specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Cadmium CD			
Analyte Program Code Challenges per Shipm			
	CD		
Beta-2-microglobulin, urine	I	3	
Cadmium, urine	I	3	
Cadmium, whole blood	I	3	
Creatinine, urine	•	3	

This program meets the Occupational Safety and Health Administration (OSHA) guidelines for proficiency testing (OSHA standard-29 CFR 1910.1027AppF).

Program Information

- Three 6.0-mL whole blood specimens and three 12.0-mL urine specimens
- Conventional and International System of Units (SI) reporting offered
- Six shipments per year

Cerebrospinal Fluid Chemistry and Oligoclonal Bands M, OLI				
Analyte	Progr	Program Code Challenges per Shipment		
	М	OLI		
Albumin, quantitative	1	ı	3	
Electrophoresis (albumin and gamma globulin)		•	3	
Glucose		ı	3	
IgG, quantitative	1	ı	3	
Lactate	1	ı	3	
Lactate dehydrogenase (LD)	1	ı	3	
Protein, total	•	ı	3	
Oligoclonal bands		ı	3	

Program Information

- M Three 5.0-mL simulated liquid spinal fluid specimens
- OLI Three 5.0-mL simulated liquid spinal fluid specimens and three 1.0-mL paired serum specimens; CSF IgG index and synthesis rate calculation challenges for each paired specimen and one online educational pattern interpretation per mailing
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year



Cystatin C CYS			
Analyte Program Code Challenges per Shipme			
CYS			
Cystatin C	ı	2	

- Two 1.0-mL liquid serum specimens
- Two shipments per year

Fecal Calprotectin FCAL			
Analyte	Program Code	Challenges per Shipment	
	FCAL		
Fecal calprotectin	I	3	

Fecal Fat FCFS				
Analyte	Program Code Challenges per Sh			
	FCFS			
Fecal fat, qualitative	1	2		

Fructosamine FT			
Analyte	Program Code Challenges per Shipme		
	FT		
Fructosamine		2	

Glucose-6-Phosphate Dehydrogenase G6PDS		
Analyte Program Code Challenges per Shipmer		
	G6PDS	
G6PD, qualitative and quantitative	I	2

H. pylori Breath Test HPBT			
Analyte Program Code Challenges per Shipme			
НРВТ			
H. pylori breath test	•	2	

Lipoprotein-Associated Phospholipase A ₂ PLA		
Analyte	Program Code	Challenges per Shipment
	PLA	
Lipoprotein-associated phospholipase (Lp-PLA ₂) activity	ı	2

- Three 1.0-g simulated fecal specimens
- Two shipments per year

Program Information

- Two 10.0-g simulated fecal fat specimens
- For microscopic detection of neutral fats (triglycerides) and/or split fats (total free fatty acids)
- Two shipments per year

Program Information

- Two 1.0-mL liquid serum specimens
- Two shipments per year

Program Information

- Two 0.5-mL lyophilized hemolysate specimens
- Two shipments per year

Program Information

- Two gas bags for qualitative reporting with the Meridian BreathID
- Two shipments per year

- Two 2.0-mL lyophilized serum specimens
- Two shipments per year

Lipoprotein Electrophoresis LPE		
Analyte/Procedure	Program Code	Challenges per Shipment
	LPE	
Lipoprotein electrophoresis	I	2

- Two 1.0-mL liquid specimens
- Two shipments per year

Protein Electrophoresis SPE, UBJP			
Analyte	Program Code Challenge		Challenges per Shipment
	SPE	UBJP	
IgA, quantitation	1		2
IgG, quantitation			2
IgM, quantitation			2
M-component (paraprotein) identification	1		2
Protein, total			2
Protein electrophoresis, serum			2
Urine Bence Jones protein			2

Program Information

- SPE Two 1.0-mL lyophilized serum specimens; one online educational protein electrophoresis challenge per mailing
- UBJP Two 10.0-mL urine specimens
- Two shipments per year



Lamellar Body Count LBC		
Procedure	Program Code	Challenges per Shipment
	LBC	
Lamellar body count	I	3

Program Information

- Three 2.0-mL simulated amniotic fluid specimens
- For use with LBC methods performed on all hematology analyzers
- Two shipments per year

Plasma Hemoglobin PHG		
Analyte Program Code Challenges per Shipmer		
	PHG	
Plasma hemoglobin		2

Program Information

- Two 2.0-mL liquid specimens
- Two shipments per year

Procalcitonin PCT		
Analyte	Program Code	Challenges per Shipment
	PCT	
Procalcitonin	ı	3

- Three 1.0-mL lyophilized serum specimens
- Two shipments per year

Pseudocholinesterase C7		
Analyte	Program Code	Challenges per Shipment
	C7	
Pseudocholinesterase		1

Salivary Cortisol SALC			
Analyte	Program Code Challenges per Shipmen		
	SALC		
Salivary cortisol	ı	3	

Accuracy-Based Testosterone, Estradiol ABS **Analyte Program Code** Challenges per Shipment **ABS** Albumin 3 Cortisol 3 Estradiol Follicle-stimulating hormone (FSH) 3 3 Luteinizing hormone (LH) 3 Prostate-specific antigen (PSA), total Sex hormone-binding globulin 3 (SHBG) 3 Testosterone ı Thyroid-stimulating hormone (TSH)

The Centers for Disease Control and Prevention (CDC) will set target values for testosterone and estradiol using the established reference methods.

Total Bile Acids TBLA		
Analyte	Program Code	Challenges per Shipment
	TBLA	
Total bile acids	I	3

Program Information

- One 2.0-mL lyophilized serum specimen
- Three shipments per year

Program Information

- Three 2.0-mL synthetic oral fluid specimens
- Two shipments per year

Program Information

- Three 1.0-mL human serum specimens
- Two shipments per year

- Three 5.0-mL liquid serum specimens
- Two shipments per year

Trace Metals R		
Analyte	Program Code	Challenges per Shipment
	R	
Aluminum		3
Chromium		3
Copper	I	3
Manganese		3
Selenium		3
Zinc		3

- Three 6.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Trace Metals, Urine TMU			
Analyte	Program Code	Challenges per Shipment	
	TMU		
Aluminum	I	3	
Arsenic	I	3	
Chromium	I	3	
Cobalt	I	3	
Copper	I	3	
Lead	I	3	
Manganese	I	3	
Mercury	I	3	
Selenium	I	3	
Thallium	I	3	
Zinc	I	3	

Program Information

- Three 12.0-mL urine specimens
- Conventional and International System of Units (SI) reporting offered
- For laboratories that monitor trace metals at normal and toxic levels
- Two shipments per year

Trace Metals, Whole Blood TMWB			
Analyte	Program Code	Challenges per Shipment	
	TMWB		
Aluminum		3	
Arsenic, total	I	3	
Chromium	I	3	
Cobalt		3	
Copper	I	3	
Manganese	I	3	
Mercury		3	
Selenium	I	3	
Thallium		3	
Zinc		3	

- Three 6.0-mL whole blood specimens
- Conventional and International System of Units (SI) reporting offered
- For laboratories that monitor trace metals at normal and toxic levels
- · Two shipments per year

Sweat Analysis Series SW2, SW4				
Analyte	alyte Program Code Challenges per Shipment			
	SW2, SW4			
Chloride	I	3		
Conductivity	I	3		

For method compatibility, see chart below.

Program Information

- SW2, SW4 Three 5.0-mL simulated liquid human sweat specimens
- Two shipments per year

Sweat Analysis Series Compatibility Matrix

Method/Procedure	Program Code		Materials Included
	SW2	SW4	
ELITechGroup and Nanoduct® Systems	•		22-gauge blunt-tipped needles
All other methodologies			No additional materials provided

Viscosity V				
Analyte Program Code Challenges per Shipmer				
V				
Viscosity		2		

Viscosity 2 Soluble Transferrin Receptor STFR

Program Information

- Two 10.0-mL serum specimens
- · Two shipments per year

Analyte Program Code Challenges per Shipment STFR Soluble transferrin receptor (sTfR) 3

Program Information

- Three 2.5-mL liquid human serum specimens
- Two shipments per year

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- Correlate results with other laboratories or instruments.
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- Utilize material with confirmed results as an alternative external quality control.
- Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Cerebrospinal Fluid, Validated Material

Validated Material	Program Code	Corresponding Program	Page
Cerebrospinal Fluid	MVM	M	74

Program Information

• Three 5.0-mL simulated liquid spinal fluid specimens

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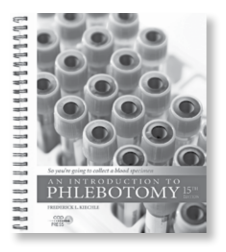
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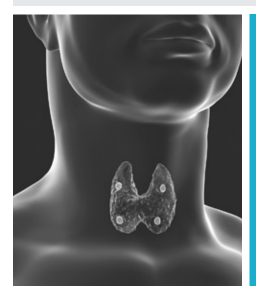
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7 Endocrinology



Be confident in the accuracy of your endocrinology testing.

Test your laboratory's proficiency with our new PT/EQA program exclusively for parathyroid hormone levels.

All current and new 2025 Centers for Medicare & Medicaid Services (CMS) regulated analytes are listed in **bold** type.

New Programs



Parathyroid Hormone (PTH)......86

Analyte Changes

CA 125 added to Ligand—General (K/KK) and removed from Tumor Markers (TM/TMX)82

Discontinued Programs

Bone Markers and Vitamins (BMV6) Bone and Mineral Metabolism, Urine (BU)

Endocrinology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Ligand-	-General K/KK	
Analyte	Program Code	Challenges per Shipment
	K/KK	
Alpha-fetoprotein (AFP)	I	5
CA 125 NEW	I	5
Carcinoembryonic antigen (CEA)	I	5
Cortisol	I	5
Ferritin	I	5
Folate, serum	I	5
Human chorionic gonadotropin (hCG), quantitative	•	5
Immunoglobulin E (IgE)	ı	5
Prostate-specific antigen (PSA), total	1	5
p2PSA	ı	5
Prostate-specific antigen, complexed (cPSA)	ı	5
Prostate-specific antigen (PSA), free	ı	5
Prostatic acid phosphatase (PAP)	I	5
Triiodothyronine (T3), free	1	5
Triiodothyronine (T3), total	1	5
T3 uptake and related tests	I	5
Thyroxine (T4), free	1	5
Thyroxine (T4), total	1	5
Thyroid-stimulating hormone (TSH)	I	5
Vitamin B ₁₂	ı	5

Program Information

- K Five 5.0-mL liquid serum specimens
- KK Five 5.0-mL liquid serum specimens in duplicate
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



MMA and Active B ₁₂ MMA			
Analyte/Procedure Program Code Challenges per Shipment			
	MMA		
Active vitamin B ₁₂	I	3	
Methylmalonic acid	I	3	

- Three 1.0-mL lyophilized serum specimens
- Two shipments per year

B-type Natriuretic Peptides BNP5			
Analyte	Program Code	Challenges per Shipment	
	BNP5		
BNP	I	5	
NT-proBNP		5	

Additional Information

- For i-STAT, Quidel Triage, and Pathfast, use Point-of-Care Cardiac Markers programs PCARM or PCARMX.
- For second instrument reporting options, see the Quality Cross Check program, BNPQ, below.

Program Information

- Five 1.0-mL liquid plasma specimens
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year

Quality Cross Check—B-type Natriuretic Peptides BNPQ			
Analyte	Program Code	Challenges per Shipment	
	BNPQ		
BNP	I	3	
NT-proBNP	ı	3	

This program does not meet regulatory requirements for proficiency testing; see program BNP5, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

- Three 1.5-mL liquid specimens
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Sex Hormones Y/YY			
Analyte	Program Code	Challenges per Shipment	
	Y/YY		
11-deoxycortisol	I	5	
17-hydroxyprogesterone	I	5	
Androstenedione	1	5	
DHEA sulfate	ı	5	
Estradiol	I	5	
Estriol, unconjugated (uE3)	1	5	
Follicle-stimulating hormone (FSH)	ı	5	
Growth hormone (GH)	I	5	
IGF-1 (somatomedin C)	1	5	
Luteinizing hormone (LH)	I	5	
Progesterone	I	5	
Prolactin	ı	5	
Sex hormone-binding globulin (SHBG)	I	5	
Testosterone	I	5	
Testosterone, bioavailable (measured)	ı	5	
Testosterone, free (measured)	I	5	

- Y Five 5.0-mL liquid serum specimens in duplicate
- YY Five 5.0-mL liquid serum specimens in triplicate
- · Conventional and International System of Units (SI) reporting offered
- Three shipments per year



Antimüllerian Hormone AMH		
Analyte	Program Code	Challenges per Shipment
	АМН	
Antimüllerian hormone		3

25-OH Vitamin D, Total VITD					
Analyte Program Code Challenges per Shipme					
VITD					
25-OH vitamin D, total ■ 3					

Program Information

- Three 1.0-mL lyophilized serum specimens
- Two shipments per year

- Three 1.0-mL liquid serum specimens
- · Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Bone and Growth BGS				
Analyte	Program Code	Challenges per Shipment		
	BGS			
IGF-1 (somatomedin C)		3		
Osteocalcin		3		

Program InformationThree 1.0-mL liquid

- Three 1.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Accuracy-Based Vitamin D ABVD					
Analyte Program Code Challenges per Shipm					
ABVD					
25-OH vitamin D (D2 and D3)	1	3			
Calcium	I	3			

Additional Information

- The Centers for Disease Control and Prevention (CDC) will establish reference targets using isotope-dilution LC-MS/MS method.
- Specimens are collected by a modified application of Clinical and Laboratory Standards Institute Guideline CLSI C37-A, Preparation and Validation of Commutable Frozen Human Serum Pools as Secondary Reference Materials for Cholesterol Measurement Procedures; Approved Guideline.

Program Information

- Three 1.0-mL liquid human serum specimens
- Serum is from multi-donor endogenous pools.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

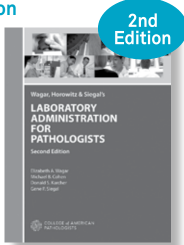
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Bone Markers and Vitamins BMV1, BMV2, BMV3, BMV4, BMV5						
Analyte		Program Code			Challenges per Shipment	
	BMV1	BMV2	BMV3	BMV4	BMV5	
1,25-dihydroxy vitamin D						3
Bone-specific alkaline phosphatase						3
Vitamin A						3
Vitamin E, total						3
C-telopeptide						3

- BMV1-4 Three 5.0-mL liquid serum specimens for each program
- BMV5 Three 1.0-mL liquid serum specimens
- Two shipments per year

Insulin, Gastrin, and C-peptide ING					
Analyte Program Code Challenges per Ship					
ING					
C-peptide		3			
Gastrin	I	3			
Insulin		3			

Insulin, Gastrin, and C-peptide ING			
Analyte	Program Code	Challenges per Shipment	
	ING		
C-peptide	I	3	
Gastrin	I	3	
nsulin	ı	3	

• Three 5.0-mL lyophilized serum specimens · Conventional and

Program Information

- International System of Units (SI) reporting offered
- Two shipments per year

Parathyroid Hormone PTH			
Analyte/Procedure	Program Code	Challenges per Shipment	
	PTH		
Parathyroid hormone (PTH)		5	

- Five 2.0-mL lyophilized serum specimens
- · Conventional and International System of Units (SI) reporting offered
- Three shipments per year

Quality Cross Check—Parathyroid Hormone PTHQ				
Analyte	Program Code Challenges per Shipment			
	PTHQ			
Parathyroid hormone (PTH)	ı	3		

This program does not meet regulatory requirements for proficiency testing; see program PTH on page 86. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

Three 2 0-ml lyonh

- Three 2.0-mL lyophilized serum specimens in duplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Accuracy-Based Glucose, Insulin, and C-peptide ABGIC						
Analyte	Program Code	Challenges per Shipment				
	ABGIC	ABGIC				
C-peptide	I	3				
Gastrin NEW	I	3				
Glucose	I	3				
Insulin		3				

Target values for C-peptide are established by isotope-dilution mass spectrometry performed at the University of Missouri, Diabetes Diagnostic Laboratory.

Second Trimester Maternal Screening FP/FPX				
Analyte Program Code Challenges per Ship				
	FP/FPX			
Alpha-fetoprotein (AFP), amniotic fluid	1	2		
Alpha-fetoprotein (AFP), serum	•	5		
Dimeric inhibin A (DIA)	I	5		
Estriol, unconjugated (uE3)	1	5		
Human chorionic gonadotropin (hCG), quantitative	ı	5		

The CAP designed these programs for laboratories using AFP and hCG for prenatal screening purposes only. For all other applications, see program K or KK on page 82.

Program Information

- Three 1.0-mL serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

- FP Five 1.0-mL serum specimens; two 1.0-mL simulated amniotic fluid specimens
- FPX All program FP serum specimens in duplicate; two 1.0-mL simulated amniotic fluid specimens
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year

First Trimester Maternal Screening FP1T, FP1B			
Analyte	Program Code Challenges per Shipmer		
	FP1T	FP1B	
Total hCG			5
Free beta hCG			5
PAPP-A			5

The CAP designed these programs for laboratories using hCG for prenatal screening purposes only. For all other applications, see program K or KK on page 82.

Program Information

- FP1T, FP1B Five 1.0-mL serum specimens
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year

Noninvasive Prenatal Testing NIPT			
Analyte	te Program Code		
Cell-free DNA screening for fetal aneuploidy	I	3	

Noninvasive prenatal testing is an exercise and is not considered proficiency testing. This exercise may be used to meet the requirements for alternative assessment.

Program Information

- · Three liquid specimens
- · Two shipments per year

Erythropoietin EPO				
Analyte	Program Code Challenges per Shipment			
EPO				
Erythropoietin	ı	2		

Fetal Fibronectin FF Analyte **Program Code** Challenges per Shipment FF 2 Fetal fibronectin

Red Blood Cell Folate FOL			
Analyte	Program Code	Challenges per Shipment	
	FOL		
RBC folate		2	

Program Information

- Two 1.5-mL serum specimens
- · Two shipments per year

Program Information

- Two 1.2-mL liquid specimens
- · Two shipments per year

- Two 2.0-mL lyophilized whole blood specimens
- · Conventional and International System of Units (SI) reporting offered
- · Three shipments per year

Renin and Aldosterone RAP					
Analyte	Program Code Challenges per Shipment				
RAP					
Aldosterone	I	3			
Renin ■ 3					

- Three 2.0-mL liquid serum specimens
- · Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Tumor Markers TM/TMX				
Analyte	Program Code	Challenges per Shipment		
	TM/TMX			
Adrenocorticotropic hormone (ACTH)	I	3		
Beta-2 microglobulin	I	3		
CA 15-3	I	3		
CA 19-9	I	3		
CA 27.29	ı	3		
CA 72-4	I	3		
Calcitonin	I	3		
Thyroglobulin	ı	3		

Program Information

- TM Three 2.0-mL liquid serum specimens
- TMX All program TM specimens in duplicate
- Two shipments per year

Human Epididymis Protein 4 HUEP		
Analyte	Program Code	Challenges per Shipment
	HUEP	
Human epididymis protein 4	1	3

- Three 1.0-mL lyophilized serum specimens
- Two shipments per year

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- Identify and troubleshoot instrument/method problems.
- Correlate results with other laboratories or instruments.
- · Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- · Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Endocrinology, Validated Materials

Validated Material	Program Code	Corresponding Program	Page
Ligand—General	KVM	K	82
Sex Hormones	YVM	Υ	84

Program Information

- KVM Five 5.0-mL liquid serum specimens; three shipments per year
- YVM Five 5.0-mL liquid serum specimens in duplicate; three shipments per year

Lead your organization in laboratory stewardship.

With immense pressure to provide fast, accurate results with limited resources, your laboratory will benefit from the CAP's **Test Ordering Program**.

Guide this effort in your organization and

- Find ways to use your resources more efficiently.
- Build your laboratory stewardship programs.
- Review your testing patterns for efficacy and utility.

The Test Ordering Program—now available to CAP customers—includes analytical tools, the latest expert-written recommendations, and suggested interventions.





8

Blood Gas, Critical Care, and Oximetry



Our programs closely mimic patient testing to ensure accuracy.

- Test specimen levels that reflect clinical decision points.
- Our reconfigured programs better meet today's blood gas laboratory needs.

All current and new 2025 Centers for Medicare & Medicaid Services (CMS) regulated analytes are listed in **bold** type.

Blood Gas, Critical Care, and Oximetry

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Critical Care Blood Gas AQ, AQH				
Analyte	Program Code		Challenges per Shipment	
	AQ	AQH		
Calcium, ionized	ı	•	2	
Chloride	ı		5	
Creatinine	I		5	
Glucose	ı		5	
Hematocrit			5	
Hemoglobin, estimated			5	
Lactate	ı	I	2	
Magnesium, ionized	ı		2	
pCO ₂	I		5	
рН	ı	I	5	
pO ₂	ı		5	
Potassium	I	•	5	
Sodium	I		5	
tCO ₂	ı	I	5	
Urea nitrogen (BUN)	I	•	5	

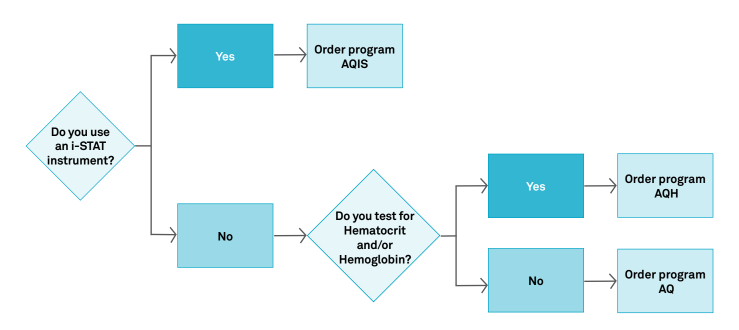
Program Information

- AQ Five 2.5-mL aqueous specimens in duplicate; appropriate for all methods except i-STAT
- AQH Five 2.5-mL aqueous specimens in duplicate and five 2.5-mL specimens for hematocrit testing in duplicate; appropriate for all methods except i-STAT
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



For multiple instrument reporting options, see the Quality Cross Check programs, AQQ and AQHQ, on page 94.

It is not appropriate to report hemoglobin and hematocrit results by co-oximetry in these programs.



Critical Care Blood Gas, i-STAT AQIS				
Analyte	Program Code	Challenges per Shipment		
	AQIS			
Calcium, ionized	1	2		
Chloride	1	5		
Creatinine	ı	5		
Glucose	1	5		
Hematocrit	1	5		
Hemoglobin, estimated	ı	5		
Lactate	1	2		
pCO ₂	1	5		
рН	ı	5		
pO ₂	1	5		
Potassium	1	5		
Sodium	ı	5		
tCO ₂	1	5		
Urea nitrogen (BUN)	•	5		

For multiple instrument reporting options, see the Quality Cross Check program, AQSQ, on page 94.

Program Information

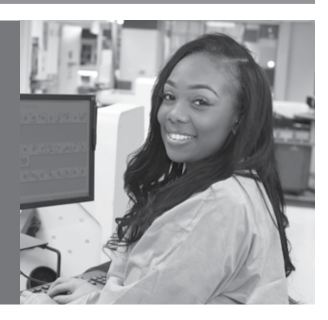
- AQIS Five specimens in duplicate for i-STAT only
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year



With direct transmission, less equals more.

Transmit your quantitative PT/EQA results directly to the CAP with direct transmission. Your laboratory will spend less time manually entering results, which will free up resources for other priorities. Plus, you will reduce clerical errors and streamline your process to be more like patient testing.

Get connected. Learn more at cap.org



Quality Cross Check—Critical Care Blood Gas AQQ, AQHQ, AQSQ				
Analyte	F	Program Cod	le	Challenges per Shipment
	AQQ	AQHQ	AQSQ	
Calcium, ionized	ı		•	3
Chloride	ı		•	3
Creatinine	ı		•	3
Glucose	1		•	3
Hematocrit			•	3
Hemoglobin, estimated			•	3
Lactate	1		•	3
Magnesium, ionized	1			3
pCO ₂	1		•	3
pH	1		•	3
pO ₂	1		•	3
Potassium	1		•	3
Sodium	1	ı	•	3
tCO ₂ (measured)			•	3
Urea nitrogen (BUN)	1		•	3

Additional Information

- It is not appropriate to report hemoglobin and hematocrit results by co-oximetry in these programs.
- These programs do no meet regulatory requirements for proficiency testing; see programs AQ, AQH, and AQIS on pages 92-93. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

Order program Yes **AQSQ** Do you use an i-STAT instrument? Order program Yes AQHQ Do you test for Hematocrit No and/or Hemoglobin? Order program No AQQ

- AQQ Three 2.5-mL specimens in triplicate; appropriate for all methods except i-STAT
- AQHQ Three 2.5-mL specimens in triplicate and three 2.5-mL specimens for hematocrit testing in triplicate; appropriate for all methods except i-STAT
- AQSQ Three 1.7-mL specimens in triplicate for i-STAT methods only
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Blood Oximetry SO				
Analyte	Program Code	Challenges per Shipment		
	S0			
Carboxyhemoglobin	1	5		
Hematocrit, estimated	1	5		
Hemoglobin, total	I	5		
Methemoglobin	I	5		
Oxyhemoglobin	I	5		

Additional Information

- This program is not compatible with Oxicom-2000, -2100, or -3000 whole blood oximeters.
- For multiple instrument reporting options, see the Quality Cross Check program, SOQ, below.

Program Information

- Five 1.8-mL stabilized human hemoglobin solution specimens
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year

Quality Cross Check—Blood Oximetry SOQ					
Analyte	nalyte Program Code Challenges per Shipment				
	SOQ				
Carboxyhemoglobin	I	3			
Hematocrit, estimated	I	3			
Hemoglobin, total	I	3			
Methemoglobin	I	3			
Oxyhemoglobin	I	3			

This program does not meet regulatory requirements for proficency testing; see program SO, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

- Three 1.2-mL liquid specimens in triplicate
- Report up to three instruments.
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Our PT/EQA resources are with you every step of the way.



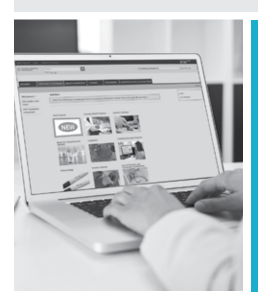
Everything you need is all in one place. With the CAP's online PT/EQA resources, you can:

- Learn the basics of the process and how to get started in our customer portal, e-LAB Solutions Suite.
- Find detailed information in our updated manual.
- Understand how to get the most out of your evaluations and participant summaries.
- Read frequently asked questions (FAQs) pertaining to performance and interpretation.

It's a great place to continue your quality journey.

Explore the CAP's online PT/EQA resources at cap.org.

9 Toxicology



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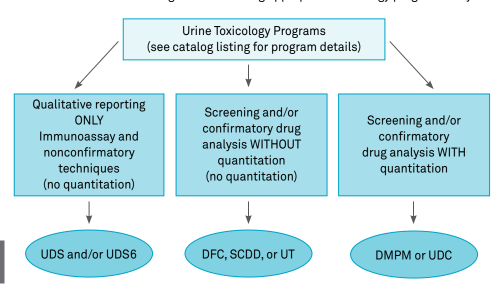
New Analyte/Drug Additions NEW

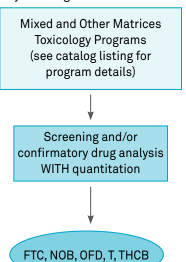
Etizolam (UT)	99
Xylazine (T, UT)	
Zolpidem carboxylic acid (DFC)	

Toxicology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Use this flowchart as a guide for ordering appropriate toxicology programs for your laboratory's testing menu.





Toxicology T			
Analyte	Program Code Challenges per Shipmer		
	Т		
See drug listing on next page	I	5	

Program Information

- A total of five specimens consisting of 20.0-mL liquid serum and 50.0-mL liquid urine specimens
- For laboratories performing qualitative and quantitative drug analysis on serum and qualitative analysis on urine specimens
- Three shipments per year

Urine Toxicology UT		
Analyte	Challenges per Shipment	
	UT	
See drug listing on next page		5

- Five 50.0-mL liquid urine specimens
- For laboratories performing qualitative drug analysis with qualitative confirmatory testing
- Three shipments per year

T and UT Programs Drug Listing

Challenges will include a mix of drugs from the list below.

6-acetylmorphine (6-AM) 7-aminoclonazepam 7-aminoflunitrazepam 7-hydroxymitragynine Acetaminophen Alpha-hydroxyalprazolam Alprazolam Amitriptyline Amphetamine Amphetamine group Aripiprazole Atenolol Atropine Barbiturate group Benzodiazepine group Benzoylecgonine Brompheniramine Buprenorphine Bupropion Butalbital Cannabinoids	Delta-9-THC (serum only) Delta-9-THC-COOH Demoxepam Desipramine Desmethylclomipramine Desmethylcyclobenzaprine* Dextromethorphan Diazepam Dihydrocodeine Diltiazem Diphenhydramine Doxepin Doxylamine Duloxetine Ecgonine methyl ester Ephedrine Etizolam NEW Fentanyl Flunitrazepam Fluoxetine Gabapentin	Meta-chlorophenylpiperazine (m-CPP) Methadone Methadone metabolite (EDDP) Methamphetamine Methylenedioxy- amphetamine (MDA) Methylenedioxy- methamphetamine (MDMA) Methylenedioxy- pyrovalerone (MDPV) Methylphenidate Metoprolol Mirtazapine Mitragynine (Kratom) Morphine N-desmethyltramadol Naproxen Norbuprenorphine Norchlordiazepoxide Norclomipramine	Nortriptyline Norverapamil O-desmethyltramadol Olanzapine Opiate group Oxazepam Oxycodone Oxymorphone Paroxetine Pentobarbital Phencyclidine Pheniramine Phenobarbital Phentermine Phenylephrine Phenyloin Pregabalin Propoxyphene Propranolol Pseudoephedrine Quetiapine
Benzoylecgonine Brompheniramine Buprenorphine Bupropion	Etizolam NEW Fentanyl Flunitrazepam Fluoxetine	Morphine N-desmethyltramadol Naproxen Norbuprenorphine Norchlordiazepoxide	Pregabalin Propoxyphene Propranolol Pseudoephedrine

^{*}Same compound

CAP/ADLM Urine Drug Testing, Screening UDS, UDS6 Analyte **Program Code** Challenges per Shipment **UDS** Limited 5 3 6-acetylmorphine (6-AM) 5 Acetaminophen 3 5 3 **Amphetamine** 5 3 Amphetamine/methamphetamine group 5 3 Barbiturate group 5 3 Benzodiazepine group Benzoylecgonine/cocaine metabolites 5 3 5 3 Buprenorphine and metabolites Cannabinoids 5 3 5 3 Ethanol 5 3 Fentanyl 5 3 Hydrocodone Lysergic acid diethylamide (LSD) 5 3 Meperidine 5 3 Meprobamate/carisoprodol 5 3 Methadone 5 3 5 Methadone metabolite (EDDP) 3 5 3 Methamphetamine 5 Methaqualone 3 5 Methylenedioxymethamphetamine (MDMA) 3 5 3 Opiate group 5 3 Oxycodone 5 3 Phencyclidine 5 3 Propoxyphene 5 3 Tramadol 5 3 Tricyclic group

- UDS Five 10.0-mL liquid urine specimens; three shipments per year
- UDS6 Three 10.0-mL liquid urine specimens; two shipments per year
- For laboratories performing drugs of abuse testing on urine specimens using immunoassay or other nonconfirmatory techniques only
- Participants will have access to the ADLM quarterly newsletter, Clinical & Forensic Toxicology News.



Urine Drug Adulterant/Integrity DAI		
Analyte	Program Code	Challenges per Shipment
	DAI	
Creatinine		3
Glutaraldehyde		3
Nitrite		3
Oxidants	1	3
рН		3
Specific gravity		3

- Three 25.0-mL urine specimens
- · Two shipments per year

Clinical Toxicology Testing: A Guide for Laboratory Professionals, Second Edition

This book is a practical guide to directing hospital toxicology laboratory operations. This edition features expanded sections on testing in the clinical setting, methodologies, and more user-friendly information on specific analytes. It provides the reader with a comprehensive view of what is needed—and expected—when offering a clinical toxicology service.

Contents include:

- Toxicology testing in the clinical setting, including new chapters on pediatric testing and chronic opioid therapy
- Toxicokinetics and methodologies, with new and expanded information on laboratory-developed tests, screening assays, targeted tests, and oral fluids and alternative matrices
- Specific analytes, including novel psychoactive substances and the use of medical cannabis
- Appendices on such useful topics as urine and serum screens, therapeutic drug monitoring, and proficiency testing

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Item number: PUB227 Softcover; 368 pages; 2020

CAP/ADLM Forensic Urine Drug Testing, Confirmatory UDC Program Code Analyte Challenges per Shipment UDC 6-acetylmorphine (6-AM) 10 Alpha-hydroxyalprazolam 10 Amphetamine 10 Benzoylecgonine 10 Buprenorphine 10 Butalbital 10 Codeine 10 Delta-9-THC-COOH 10 10 Fentanyl 10 Hydrocodone Hydromorphone 10 10 Lorazepam 10 Methadone Methadone metabolite (EDDP) 10 Methamphetamine 10 Methaqualone 10 Methylenedioxyamphetamine (MDA) 10 Methylenedioxyethylamphetamine 10 (MDEA) Methylenedioxymethamphetamine 10 (MDMA) Morphine 10 Norbuprenorphine ı 10 10 Nordiazepam Norfentanyl 10 Norpropoxyphene 10 10 Oxazepam 10 Oxycodone Oxymorphone 10 Phencyclidine 10 Phenobarbital 10 ı Propoxyphene 10

ı

Ī

10 10

10

10 10

Program Information

- Ten 50.0-mL liquid urine specimens
- For laboratories that perform both screening and confirmatory testing, including quantitation, for drugs of abuse in urine specimens; laboratories are asked to report creatinine, pH, and specific gravity for each specimen to ensure specimen adulteration has not occurred
- Participants will have access to the ADLM quarterly newsletter, Clinical & Forensic Toxicology News.
- · Four shipments per year



Adulterant/Integrity Indicator

Secobarbital

Temazepam

Creatinine

Specific gravity

рΗ

Analyte	Program Code OFD	Challenges per Shipment
	OED	
	OFD	
Amphetamine Group	I	5
Amphetamine	ı	5
Methamphetamine	ı	5
Methylenedioxyamphetamine (MDA)	ı	5
Methylenedioxymethamphetamine (MDMA)	ı	5
Benzodiazepine Group	I	5
Alprazolam	ı	5
Diazepam	ı	5
Nordiazepam	I	5
Oxazepam	ı	5
Temazepam	ı	5
Buprenorphine	I	5
Buprenorphine and norbuprenorphine	ı	5
Cocaine and/or metabolite	ı	5
Benzoylecgonine	ı	5
Cocaine	ı	5
Cannabinoid	ı	5
Delta-9-THC	I	5
Cotinine	I	5
Fentanyl and/or metabolite	ı	5
Fentanyl	I	5
Norfentanyl	ı	5
Methadone	ı	5
Opiate Group	I	5
6-acetylmorphine (6-AM)	ı	5
Codeine	ı	5
Hydrocodone	ı	5
Hydromorphone	ı	5
Morphine	ı	5
Oxycodone	ı	5
Oxymorphone	I	5
Phencyclidine (PCP)	ı	5

- Five 2.0-mL oral fluid specimens
- For laboratories performing drug screening, confirmation, and quantitation
- Four shipments per year



Vitreous Fluid, Postmortem VF			
Analyte	Program Code	Challenges per Shipment	
	VF		
Acetone	I	3	
Chloride	I	3	
Creatinine	I	3	
Ethanol	I	3	
Glucose	I	3	
Potassium	I	3	
Sodium	I	3	
Vitreous urea nitrogen	I	3	

- Three 5.0-mL synthetic vitreous fluid specimens
- For forensic and other toxicology laboratories that perform quantitative analysis of vitreous fluid
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year



Serum Drug Screening SDS			
Analyte	Program Code	Challenges per Shipment	
	SDS		
Acetaminophen, quantitative	I	3	
Acetone, semiquantitative and qualitative	I	3	
Barbiturate group, qualitative	I	3	
Benzodiazepine group, qualitative	I	3	
Salicylate, quantitative	I	3	
Total tricyclic antidepressants, qualitative	ı	3	

This program does not meet the regulatory requirements for proficiency testing for laboratories that quantitate Acetaminophen and/or Salicylate for TDM purposes.

Program Information
• Three 2.0-mL serum
specimens
- For laboratorios that

- For laboratories that perform serum drug screening using immunoassay or other screening techniques
- · Two shipments per year

CAP/ADLM Alcohol/Volatiles AL1, AL2			
Analyte	Progra	m Code	Challenges per Shipment
	AL1 Whole Blood	AL2 Serum	
Acetone, quantitative	•		5
Ethanol, quantitative	ı		5
Ethylene glycol, qualitative and quantitative	ı		5
Isopropanol, quantitative	I		5
Methanol, quantitative	ı		5

- AL1 Five 5.0-mL liquid whole blood specimens; conventional reporting
- AL2 Five 2.0-mL liquid serum specimens; conventional and International System of Units (SI) reporting offered
- Three shipments per year



Ethanol Biomarkers ETB		
Analyte Program Code Challenges per Si		
	ETB	
Ethyl glucuronide (EtG), qualitative and quantitative	ı	3
Ethyl sulfate (EtS), quantitative	ı	3

- Three 10.0-mL synthetic urine specimens
- Two shipments per year

CAP/ADLM Blood Lead BL				
Analyte Program Code Challenges per Shipmen				
BL				
Lead	I	5		

This program meets the Occupational Safety and Health Administration (OSHA) requirements for proficiency testing [OSHA lead standards-29 CFR 1910.1025(j)(2)(iii)].

Program Information

- Five 6.0-mL liquid nonhuman whole blood specimens
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



Cadmium CD			
Analyte	Program Code	Challenges per Shipment	
	CD		
Beta-2-microglobulin, urine	I	3	
Cadmium, urine	1	3	
Cadmium, whole blood	I	3	
Creatinine, urine	I	3	

This program meets the Occupational Safety and Health Administration (OSHA) guidelines for proficiency testing (OSHA standard-29 CFR 1910.1027AppF).

Program Information

- Three 6.0-mL whole blood specimens and three 12.0-mL urine specimens
- Conventional and International System of Units (SI) reporting offered
- Six shipments per year

Nicotine and Tobacco Alkaloids NTA				
Analyte	Program Code	Challenges per Shipment		
	NTA			
Cotinine	1	3		
Nicotine		3		

- Three 25.0-mL urine specimens
- Designed for laboratories that qualitatively and/ or quantitatively test for cotinine and/or nicotine in urine
- Two shipments per year

Trace Metals R				
Analyte	Program Code	Challenges per Shipment		
	R			
Aluminum	I	3		
Chromium	I	3		
Copper	1	3		
Manganese	1	3		
Selenium	1	3		
Zinc	ı	3		

- Three 6.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Trace Metals, Urine TMU				
Analyte	Program Code	Challenges per Shipment		
	TMU			
Aluminum	I	3		
Arsenic	I	3		
Chromium	I	3		
Cobalt	I	3		
Copper	I	3		
Lead	I	3		
Manganese		3		
Mercury	•	3		
Selenium	I	3		
Thallium	ı	3		
Zinc	ı	3		

Program Information

- Three 12.0-mL urine specimens
- Conventional and International System of Units (SI) reporting offered
- For laboratories that monitor trace metals at normal and toxic levels
- Two shipments per year

Trace Metals, Whole Blood TMWB				
Analyte	Program Code	Challenges per Shipment		
	TMWB			
Aluminum		3		
Arsenic, total		3		
Chromium		3		
Cobalt		3		
Copper		3		
Manganese		3		
Mercury		3		
Selenium		3		
Thallium		3		
Zinc		3		

- Three 6.0-mL whole blood specimens
- Conventional and International System of Units (SI) reporting offered
- For laboratories that monitor trace metals at normal and toxic levels
- Two shipments per year

Forensic Toxicology, Criminalistics FTC				
Analyte	alyte Program Code Challenges per Shipmen			
FTC				
See drug listing below	I	5		

- Five 20.0-mL whole blood specimens
- For crime and hospital laboratories that have forensic toxicology divisions performing qualitative and quantitative analysis of drugs in whole blood specimens
- Three shipments per year

FTC Program Drug Listing

Challenges will include a mix of drugs from the list below.

6-acetylmorphine (6-AM)	Desmethylsertraline	Methylenedioxyamphetamine	Oxymorphone
7-aminoclonazepam	Dextromethorphan	(MDA)	Paroxetine
7-aminoflunitrazepam	Diazepam	Methylenedioxymethamphetamine (MDMA)	Pentobarbital
7-hydroxymitragynine	Dihydrocodeine	Methylenedioxypyrovalerone	Phencyclidine
Acetaminophen	Diltiazem	(MDPV)	Phenethylamine
Alpha-hydroxyalprazolam	Diphenhydramine	Methylphenidate	Pheniramine
Alprazolam	Doxepin	Metoprolol	Phenobarbital
Amitriptyline	Doxylamine	Midazolam	Phentermine
Amphetamine	Duloxetine	Mirtazapine	Phenylephrine
Aripiprazole	Ecgonine ethyl ester	Mitragynine (Kratom)	Phenytoin
Atenolol	Ecgonine methyl ester	Morphine*	Pregabalin
Atropine	Ephedrine	N-desmethyltramadol	Propoxyphene
Benzoylecgonine	Fentanyl*	Naproxen	Propranolol
Brompheniramine	Flunitrazepam	Norbuprenorphine	Pseudoephedrine
Buprenorphine	Fluoxetine	Norchlordiazepoxide	Quetiapine
Bupropion	Gabapentin	Norclomipramine	Quinine
Butalbital	Gamma-hydroxybutyrate (GHB)	Norcodeine	Ranitidine
Carbamazepine	Hydrocodone	Norcyclobenzaprine	Ritalinic acid
Carbamazepine-10,	Hydromorphone	Nordiazepam	Salicylate
11-epoxide	Hydroxybupropion	Nordoxepin	Sertraline
Carisoprodol	Hydroxyzine	Norfentanyl	Strychnine
Chlordiazepoxide	Ibuprofen	Norfluoxetine	Tapentadol
Chlorpheniramine	Imipramine	Norketamine	Temazepam
Citalopram	Ketamine	Normeperidine	Topiramate
Clomipramine	Lamotrigine	Normirtazapine	Tramadol
Clonazepam	Levetiracetam	Noroxycodone	Trazodone
Clozapine	Lidocaine	Norpropoxyphene	Trimipramine
Cocaethylene	Lorazepam	Norsertraline	Valproic acid
Cocaine	Lysergic acid diethylamide (LSD)	Nortrimipramine	Venlafaxine
Codeine	Meperidine*	Nortriptyline	Verapamil
Cyclobenzaprine*	Mephedrone	• •	Zolpidem
Delta-9-THC	Meprobamate	Norverapamil O-desmethyltramadol	·
Delta-9-THC-COOH	Methadone		
Demoxepam	Methadone metabolite (EDDP)	Olanzapine	
Desipramine	Methamphetamine	Oxazepam	
Desmethylclomipramine	r	Oxycodone	*and/or metabolite(s

Synthetic Cannabinoid/Designer Drugs SCDD					
Analyte Program Code Challenges per Shipment					
SCDD					
Synthetic cannabinoid/designer drugs					

Synthetic cannabinoids and designer drug stimulants are widespread and constantly changing in respect to the available chemical moieties. In order to stay contemporary, the CAP has decided to modify the compounds in this program in accordance with the appearance and prevalence of new compounds.

Program Information

- Three 10.0-mL urine specimens
- For laboratories that perform screening and confirmatory testing for the compounds found in this program
- · Two shipments per year



SCDD Program Drug Listing

Challenges will include a mix of drugs.

For the most current list of drugs, please go to cap.org. Under the Laboratory Improvement tab, click on Catalog and Ordering Information. The list is located under the PT Order Supplements header.

Novel Opioids and Benzodiazepines NOB			
Analyte Program Code Challenges per Shipme			
	NOB		
Novel opioids and benzodiazepines	ı	3	

Program Information

- Three 15.0-mL whole blood specimens
- For forensic and toxicology laboratories that perform qualitative and/ or quantitative analysis of synthetic opioids and benzodiazepines
- · Two shipments per year



NOB Program Drug Listing

Challenges will include a mix of drugs.

For the most current list of drugs, please go to cap.org. Under the Laboratory Improvement tab, click on Catalog and Ordering Information. The list is located under the PT Order Supplements header.

Blood Cannabinoids THCB			
Analyte	Program Code	Challenges per Shipment	
	ТНСВ		
Delta-8-THC	I	3	
Delta-9-THC	I	3	
Delta-9-THC-COOH ■ 3			
11-hydroxy-THC ■ 3			

- Three 10.0-mL whole blood specimens
- For toxicology laboratories that perform qualitative and/or quantitative analysis of cannabinoids in blood
- Two shipments per year

Antifungal Drugs Monitoring		AFD
Analyte	Program Code	Challenges per Shipment
	AFD	
Fluconazole	I	3
Itraconazole	I	3
Posaconazole		3
Voriconazole	I	3

Program Information

- Three 2.0-mL serum specimens
- For laboratories performing quantitative analysis of antifungal agents
- · Two shipments per year

Sample Exchange Registry for Alternative Assessment

When no formal proficiency testing is yet available, join the CAP's Sample Exchange Registry. After at least three laboratories are identified as testing for the same rare analyte, the CAP can anonymously deliver a sample from each laboratory to another participating facility, all of whom then report their results to us. We send each participant a custom result report, including an anonymous participant summary covering all the laboratories that took part.

Learn more at cap.org

Drug Monitoring for Pain Management DMPM				
Analyte	Program Code	Challenges per Shipment		
DMPM				
See drug listing below		3		

- Three 40.0-mL urine specimens
- For laboratories offering qualitative, confirmatory, and/or quantitative urine drug analysis for pain management
- Includes clinical cases and questions along with detailed descriptions of how to interpret test results
- Two shipments per year

DMPM Program Drug Listing

Challenges will include a mix of drugs from the list below.

Amphetamine group	Fentanyl	Nordiazepam
6-acetylmorphine (6-AM)	Fentanyl and/or metabolites	Norfentanyl
7-aminoclonazepam	Gabapentin	Norhydrocodone
Alpha-hydroxyalprazolam	Hydrocodone	Normeperidine
Alprazolam	Hydromorphone	Noroxycodone
Amphetamine	I-amphetamine	Noroxymorphone
Barbiturate group	I-methamphetamine	Norpropoxyphene
Benzodiazepine group	Lorazepam	O-desmethyltramadol
Benzoylecgonine	Meperidine	Opiate group
Buprenorphine	Meperidine and/or metabolites	Oxazepam
Buprenorphine and/or metabolites	Meprobamate	Oxycodone
Butalbital	Methadone	Oxymorphone
Cannabinoids	Methadone metabolite (EDDP)	Phenobarbital
Carisoprodol	Methamphetamine	Pregabalin
Carisoprodol and/or metabolites	Methylenedioxyamphetamine (MDA)	Propoxyphene
Clonazepam	Methylenedioxymethamphetamine	Propoxyphene and/or metabolites
Cocaine	(MDMA)	Tapentadol
Cocaine and/or metabolites	Morphine	Tapentadol-O-sulfate
Codeine	N-desmethyltramadol	Temazepam
Delta-9-THC-COOH	Naloxone	Tramadol
Diazepam	Norbuprenorphine	Tramadol and/or metabolites

Drug-Facilitated Crime DFC			
Analyte	Program Code	Challenges per Shipment	
	DFC		
See drug listing below		3	

- Three 25.0-mL urine specimens
- For laboratories performing qualitative urine drug analysis with confirmation testing
- Designed for laboratories performing testing for drugs associated with drugfacilitated crimes, which target drugs at much lower concentrations than in other toxicology programs
- Two shipments per year



DFC Program Drug Listing

Challenges will include a mix of drugs from the list below.

4-hydroxytriazolam	Gabapentin	Norvenlafaxine
7-aminoclonazepam	Gamma hydroxybutyrate (GHB)	O-desmethyltramadol
7-aminoflunitazepam	Hydrocodone	Oxazepam
Alpha-hydroxyalprazolam	Hydromorphone	Oxycodone
Amitriptyline	Hydroxyzine	Oxymorphone
Amobarbital	Imipramine	Paroxetine
Amphetamine	Ketamine	Pentobarbital
Benzoylecgonine	Lorazepam	Phencyclidine (PCP)
Bromazepam	Meperidine	Phenobarbital
Brompheniramine	Meprobamate	Phenytoin
Butalbital	Meta-chlorophenylpiperazine	Promethazine
Carisoprodol	(m-CPP)	Propoxyphene
Chlorpheniramine	Methadone	Quetiapine
Citalopram/escitalopram	Methadone metabolite (EDDP)	Scopolamine
Clobazam	Methamphetamine	Secobarbital
Clonidine	Methylenedioxyamphetamine (MDA)	Sertraline
Clozapine	Methylenedioxymethamphetamine (MDMA)	Tapentadol
Codeine	(MDMA) Midazolam	Temazepam
Cyclobenzaprine		Tetrahydrozoline
Delta-9-THC-COOH	Morphine	Topiramate
Desipramine	Norbuprenorphine	Tramadol
Dextromethorphan	Nordoxepin	Valproic acid
Diphenhydramine	Norfentanyl	Venlafaxine
Doxepin	Norfluoxetine	Zaleplon
Doxylamine	Norketamine	Ziprasidone
Estazolam	Normeperidine	Zolpidem
Ftizolam	Norpropoxyphene	Zolpidem carboxylic aci

Etizolam Zolpidem carboxylic acid NEW Fentanyl Zopiclone/Eszopiclone Nortriptyline

Fluoxetine

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- Identify and troubleshoot instrument/method problems.
- · Correlate results with other laboratories or instruments.
- Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Toxicology, Validated Material

Validated Material	Program Code	Corresponding Program	Page
Urine Drug Testing, Screening	UDSM	UDS	100

Program Information

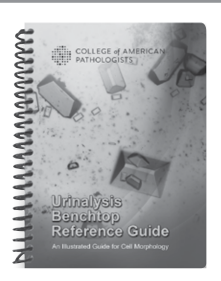
- Five 10.0-mL liquid urine specimens
- Three shipments per year

Urinalysis Benchtop Reference Guide

- Thirty-four different cell identifications, including common and rare cells
- Detailed descriptions for each cell morphology
- Eight tabbed sections for easy reference
- A durable and water-resistant format to withstand years of benchtop use—5" x 6½"

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Item number: UABRG

Spiral bound; 38 pages; 2014



The CAP's Accuracy-Based Programs do what proficiency testing can't.

- Use the CAP's Accuracy-Based Programs to verify the accuracy of your test results against a gold standard.
- Accuracy-Based Programs use challenge specimens that are matrix-related, bias-free, and have target values traceable to certified reference materials.
- Only the CAP's Accuracy-Based Programs allow laboratories to compare their test results with reference method results.

Accuracy-Based Programs

Accuracy-Based ProgramsValidated Materials	
New Analyte Additions NEW	
Accuracy-Based Glucose, Insulin and C-peptide (ABGIC)	118

Accuracy-Based Programs

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Accuracy-Based Lipids ABL			
Analyte	Program Code	Challenges per Shipment	
	ABL		
Apolipoprotein A1	I	3	
Apolipoprotein B	I	3	
Cholesterol*	I	3	
HDL cholesterol*	I	3	
Non-HDL cholesterol	I	3	
LDL cholesterol	I	3	
Lipoprotein(a)	I	3	
Triglycerides*	I	3	

^{*}This analyte will be evaluated against the reference method.

Program Information

- Three 1.0-mL human serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Accuracy-Based Vitamin D ABVD			
Analyte	Program Code	Challenges per Shipment	
	ABVD		
25-OH vitamin D (D2 and D3)	•	3	
Calcium	I	3	

Additional Information

- The Centers for Disease Control and Prevention (CDC) will establish reference targets using isotope-dilution LC-MS/MS method.
- · Specimens are collected by a modified application of Clinical and Laboratory Standards Institute Guideline CLSI C37-A, Preparation and Validation of Commutable Frozen Human Serum Pools as Secondary Reference Materials for Cholesterol Measurement Procedures; Approved Guideline.

Program Information

- Three 1.0-mL liquid human serum specimens
- Serum is from multi-donor endogenous pools.
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Accuracy-Based Testosterone, Estradiol ABS			
Analyte	Program Code	Challenges per Shipment	
	ABS		
Albumin	ı	3	
Cortisol		3	
Estradiol	•	3	
Follicle-stimulating hormone (FSH)	I	3	
Luteinizing hormone (LH)	•	3	
Prostate-specific antigen (PSA), total	I	3	
Sex hormone-binding globulin (SHBG)	ı	3	
Testosterone		3	
Thyroid-stimulating hormone (TSH)		3	

The Centers for Disease Control and Prevention (CDC) will set target values for testosterone and estradiol using the established reference methods.

Accuracy-Based Urine ABU			
Analyte	Program Code	Challenges per Shipment	
	ABU		
Calcium		3	
Creatinine		3	
Protein, total		3	
Urine albumin, quantitative		3	
Urine albumin: creatinine ratio		3	

Program Information

- Three 1.0-mL human serum specimens
- Two shipments per year

Program Information

- Three 5.0-mL human urine specimens
- Two shipments per year

Creatinine Accuracy Calibration Verification/Linearity LN24			
Analyte	Program Code		
	LN24	LN24 Target Range	
Creatinine	I	0.6-4.0 mg/dL	
Estimated glomerular filtration rate (eGFR)	ı		

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

eGFR results will be evaluated with a calculation verification comparison.

The College of American Pathologists (CAP) and the National Kidney Disease Education Program (NKDEP) have an initiative to harmonize clinically reported creatinine values. This initiative is analogous to what the federal health agencies and the clinical laboratory community did to improve the accuracy of cholesterol and glycohemoglobin testing.

Program Information

- Six 1.0-mL human serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Harmonized Thyroid ABTH					
Analyte	nalyte Program Code Challenges per Shipme				
	ABTH				
Triiodothyronine (T3), free	I	3			
Triiodothyronine (T3), total	I	3			
Thyroxine (T4), free	I	3			
Thyroxine (T4), total	I	3			
Thyroid-stimulating hormone (TSH)	I	3			

Additional Information

- Analytes will be evaluated using harmonization.
- Specimens are collected by a modified application of Clinical and Laboratory Standards Institute Guideline CLSI C37-A, Preparation and Validation of Commutable Frozen Human Serum Pools as Secondary Reference Materials for Cholesterol Measurement Procedures; Approved Guideline.

Program Information

- Three 1.0-mL frozen human serum specimens
- Two shipments per year

Hemoglobin A1c Accuracy Calibration Verification/Linearity LN15			
Analyte	Program Code		
	LN15	LN15 Target Range	
Hemoglobin A1c	I	5%-12%	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

CAP-assigned target values are derived from Hemoglobin A1c measurements assayed by National Glycohemoglobin Standardization Program (NGSP) secondary reference laboratories.

Hemoglobin A1c Waived GH2		
Analyte	Program Code	Challenges per Shipment
	GH2	
Hemoglobin A1c		3

Additional Information

- This program will be evaluated against the National Glycohemoglobin Standardization Program (NGSP) reference method.
- For multiple instrument reporting options, see the Quality Cross Check program, GHQ, see page 62.
- This program has limited stability. Laboratories outside the US or Canada should consider purchasing GH5I, which has longer stability.

Hemoglobin A1c GH5		
	Program Code	Challenges per Shipment
	GH5	
Hemoglobin A1c		5

Additional Information

- This program will be evaluated against the National Glycohemoglobin Standardization Program (NGSP) reference method.
- For multiple instrument reporting options, see the Quality Cross Check program, GHQ, see page 62.
- This program has limited stability. Laboratories outside the US or Canada should

Program Information

- Six 0.8-mL liquid human whole blood specimens
- · Two shipments per year

Program Information

- · Accuracy-Based program
- Three 0.8-mL liquid human whole blood specimens
- Two shipments per year
- · Designed for waived methods

Program Information

- Accuracy-Based program
- Five 0.8-mL liquid human whole blood specimens
- · Three shipments per year

10

Accuracy-Based Glucose, Insulin, and C-peptide ABGIC			
Analyte	Program Code	Challenges per Shipment	
	ABGIC		
C-peptide	I	3	
Gastrin NEW	I	3	
Glucose	I	3	
Insulin	I	3	

Target values for C-peptide are established by isotope-dilution mass spectrometry performed at the University of Missouri, Diabetes Diagnostic Laboratory.

Program Information

- Three 1.0-mL serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Color Atlas of Hematology—Vol 1. Peripheral Blood Color Atlas of Hematology—Vol 2. Bone Marrow

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Vol 1. Peripheral Blood

Item number: PUB222 Hardcover; 480 pages; 2018

Vol 2. Bone Marrow

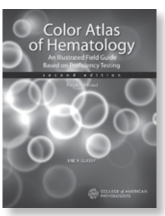
Item number: PUB229 Hardcover; 408 pages; 2023

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Validated Materials

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- Identify and troubleshoot instrument/method problems.
- Correlate results with other laboratories or instruments.
- Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

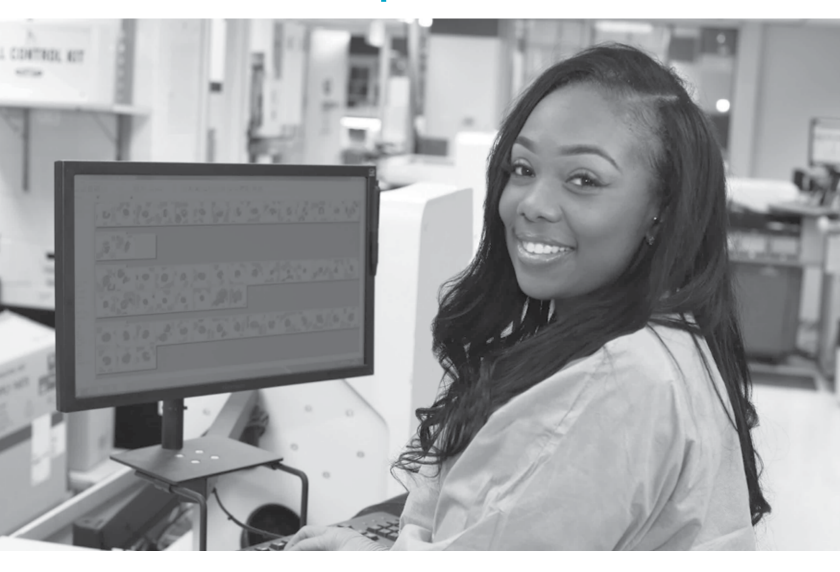
Chemistry, Validated Materials			
Validated Material	Validated Material Code	Corresponding Program	Page
General Chemistry and Therapeutic Drugs	CZVM	CZ	54-56
Cerebrospinal Fluid	MVM	M	74
Urine Chemistry—General	UVM	U	68

Coagulation—Limited, Validated Material				
Validated Material	Validated Material Code	Corresponding Program	Page	
Coagulation—Limited	CGM	CGL	166	

Endocrinology, Validated Materials			
Validated Material Code Corresponding Program Pag			
Ligand—General KVM K			
Sex Hormones	YVM	Υ	84

Toxicology, Validated Material			
Validated Material Code Corresponding Program Page			
Urine Drug Testing, Screening	UDSM	UDS	100

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Transmit your quantitative PT/EQA results directly to the CAP with direct transmission. Your laboratory will spend less time manually entering results, which will free up resources for other priorities. Plus, you will reduce clerical errors and streamline your process to be more like patient testing.

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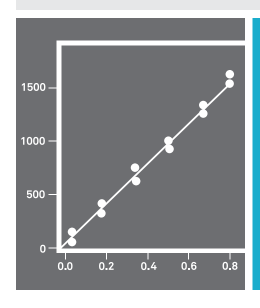
More accurate

Less time entering results

More time for patient testing



1 Instrumentation Verification Tools



Ensure your instrument and method are performing to their optimal levels.

Verify your analytical measurement range using our newest calibration verification/linearity programs for:

- Thyroid panel (LN50)
- Factor VIII (LN51)
- Hepatitis B viral load (LN52)

Instrumentation Verification Tools

Calibration Verification/Linearity	122
Instrumentation Quality Management Programs	137
New Programs NEW	
Factor VIII Calibration Verification/Linearity (LN51)	133
HBV Viral Load Calibration Verification/Linearity (LN52)	133
Thyroid Panel Calibration Verification/Linearity (LN50)	136

Discontinued Programs

Reticulocyte Calibration Verification/Linearity (LN18)

Calibration Verification/Linearity

The CAP CVL Program

The CAP is your trusted calibration verification and linearity (CVL) partner. Our CVL program will help you meet both CLIA regulations and CAP Laboratory Accreditation Program requirements for calibration and analytical measurement range verification under 42 CFR 493.1255(bX3). Don't let instrument problems impact your patient results; use the calibration verification and linearity studies to ensure your instrument and method are performing to their optimal levels.

With your enrollment in the CAP CVL program you'll receive:

· Testing Kit

- Kit Instructions—Contain important information to help you complete testing and accurately report your results
- Specimens—The majority of CAP CVL programs offer human-based materials to closely mimic your patient results

Customized Report Package

- Executive Summary—A quick overview of both your calibration verification and linearity results for all reported analytes
- Calibration Verification Evaluation
- Linearity Evaluation
 - Rapid result turnaround is complimentary for most CVL programs. View your expedited linearity evaluations within two business days of submission by logging into e-LAB Solutions Suite.
- Linearity Troubleshooting Report
- Participant Summary—A summary of laboratory performance that includes peer group statistics and enhanced diagnostic information for early insight into potential problems

Additional Tools

- Calibration Verification/Linearity Program User's Guide—Get assistance in interpreting your evaluations and reports as well as helpful troubleshooting information with suggested actions. Also available online by logging into e-LAB Solutions Suite
- Calibration Verification Troubleshooting Guide—The guide provides suggested actions if you receive a
 calibration verification result of Different, or if your evaluation result is Verified over a range that does not
 include all of your reported results.
- Calibration Verification/Linearity Surveys Investigation Checklist for Problematic Results—Interpretative checklists are included to help with troubleshooting and documentation.

Your Total Calibration Verification/Linearity (CVL) Solution			
CVL Program	Page No.	Corresponding Proficiency Testing Program	Page No.
LN2 - Chemistry, Lipid, Enzyme CVL	124	C1, C3/C3X, C4,	
LN2BV - Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124	CZ/CZX/CZ2X	54-56
LN3 - Therapeutic Drug Monitoring CVL	125	CZ/CZX/CZ2X/Z	54-56
LN5 - Ligand CVL	125		
LN5S - Ligand CVL – all Siemens ADVIA (Centaur, CP, and XP) and Atellica IM	125	K/KK	82
LN6 - Urine Chemistry CVL	126	U	68
LN7 - Immunology CVL	126	IG/IGX	216
LN8 - Reproductive Endocrinology CVL	127	Y/YY	84
LN9 - Hematology CVL	127	FH series, HE	140
LN11 - Serum Ethanol CVL	127	AL2	104
LN12 - C-reactive Protein CVL	128	CRP	216
LN13, LN13C - Blood Gas/Critical Care CVL	128	AQ, AQH, AQIS	92-93
LN15 - Hemoglobin A1c Accuracy CVL	128	GH2, GH5	62
LN16 - Homocysteine CVL	129	HMS	63
LN17 - Whole Blood Glucose CVL	129	N/A	
LN19 - Reticulocyte CVL	129	RT3	146
LN20 - Urine Albumin CVL	129	U	68
LN21 - High-Sensitivity C-reactive Protein CVL	130	HSCRP	63
LN22 - Flow Cytometry CVL	130	FL	224
LN23 - Prostate-Specific Antigen CVL	130	K/KK	82
LN24 - Creatinine Accuracy CVL	131	C1, C3/C3X, C4, CZ/CZX/CZ2X	54-56
LN25 - Troponin I CVL	131	CRT, CRTI	60
LN30 - B-type Natriuretic Peptides CVL	131	BNP5	59
LN31 - Immunosuppressive Drugs CVL	132	CS	58
LN32 - Ammonia CVL	132	C1, C3/C3X, CZ/CZX/CZ2X	54-56
LN33 - Serum Myoglobin CVL	132	CRT, CRTI	60
LN34 - Tumor Markers CVL	132	TM/TMX	89
LN35 - Thrombophilia CVL	133	CGS2	168
LN36 - Heparin CVL	133	CGS4	168
LN37 - von Willebrand Factor Antigen CVL		CGS3	168
LN38 - CMV Viral Load CVL	133 133	VLS, VLS2	206
LN39 - HIV Viral Load CVL	133		
LN40 - Vitamin D CVL	134	HIVG, HV2 VITD	206 84
LN41 - Procalcitonin CVL	134	PCT	
LN42 - D-dimer CVL	134	CGL, CGDF	76 166
LN44 - Fibrinogen CVL	134	CGL, CGDF	166
LN45 - HCV Viral Load CVL		HCV2	
LN46 - C-peptide/Insulin CVL	133 135	ING	205
LN47 - High-Sensitivity Troponin T CVL	135		86
LN48 - High-Sensitivity Troponin I CVL	135	HCRT, HCRTI	60
LN49 - Cystatin C CVL		HCRT, HCRTI	60
LN50 - Thyroid Panel CVL NEW	135	CYS	74
	136	C1, C3/C3X, CZ/CZX/CZ2X, K/KK	54-56, 8
LN51 - Factor VIII CVL NEW	133	CGE/CGEX, CGS3, ECF	167-168
LN52 - HBV Viral Load CVL NEW	133	HBVL/HBVL5	205

All CVL programs provide individual evaluation reports by analytes, an executive summary, and graphical plots for linearity and calibration verification.

11

Instrumentation Verification Tools

Chemistry, Lipid, Enzyme Calibration Verification/Linearity LN2, LN2BV Program LN2 Target Analyte **LN2BV Target Ranges** Units Code Ranges LN2, (All All Beckman Vitros LN2BV Instruments) (Except AU) ı Albumin 1.5 - 9.0g/dL Calcium 4.0 - 18.0mg/dL Chloride 60-180 mmol/L CO_2 ı 7-42 mmol/L Creatinine 0.8 - 34.0mg/dL ı 20-750 Glucose ı mg/dL 10-950 Iron ı µg/dL Magnesium ı 0.5 - 9.0mg/dL Osmolality 200-600 m0sm/kg H₂0 ı 0.5 - 22.0**Phosphorus** mg/dL Potassium 1.5 - 13.0mmol/L Protein 1.5 - 12.0g/dL Sodium 65 - 195mmol/L Urea nitrogen/Urea 5-170 mg/dL Uric acid ı 1-25 mg/dL Alkaline phosphatase ı 25-1,000 25-1,100 U/L 25-1,800 30-700 U/L ALT (SGPT) 10-900 10-650 30-900 U/L Amylase 30 - 1,80030-800 U/L AST (SGOT) ı 10-500 10-700 10-900 25-700 U/L Creatine kinase ı 25-2,000 25-1,200 1-200 CK-2 (MB) mass ı 1-250 1-300 ng/mL Gamma glutamyl ı 10-1,400 10-900 10-1,100 U/L transferase Lactate 50-700 185-3,000 U/L 50-1,800 dehydrogenase 20-190 U/L 20-1,200 150-2,500 Lipase 0.1 - 10.0Bilirubin, direct mg/dL Bilirubin, total 0.2 - 25.0mg/dL Cholesterol 35-625 mg/dL HDL ı 7-120 mg/dL 20-700 Triglycerides mg/dL

Program Information

- Seven 5.0-mL liquid serum specimens for basic chemistry, six 3.0-mL liquid serum specimens for direct and total bilirubin, seven 2.0-mL liquid serum specimens for lipids, and seven 5.0-mL liquid serum specimens for enzymes
- LN2 Appropriate for most major instruments
- LN2BV Appropriate for Beckman (except AU) and Vitros instruments only
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Therapeutic Drug Monitoring Calibration Verification/Linearity LN3		
Analyte	Program Code	
	LN3	LN3 Target Ranges
Acetaminophen	ı	20-350 μg/mL
Amikacin	I	2-45 μg/mL
Carbamazepine	I	2-25 μg/mL
Digoxin	I	0.5-4.4 ng/mL
Gentamicin	I	1–11 μg/mL
Lidocaine	I	1–10 μg/mL
Lithium	I	0.3-4.0 mmol/L
Phenobarbital	ı	8-80 μg/mL
Phenytoin	ı	5-35 μg/mL
Salicylate	ı	7-90 mg/dL
Theophylline	ı	5-35 μg/mL
Tobramycin	1	1–10 μg/mL
Valproic acid	ı	15–140 μg/mL
Vancomycin	1	7–85 μg/mL

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 4.0-mL liquid serum specimens
- A seventh 4.0-mL liquid serum specimen for acetaminophen, carbamazepine, and vancomycin
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Ligand Calibration Verification/Linearity LN5, LN5S			
Analyte	Program Code		
	LN5, LN5S*	LN5 Target Ranges	LN5S Target Ranges
AFP		1.0-900	.0 ng/mL
CEA	1	0.5-750.0 ng/mL	0.6-90.0 ng/mL
Cortisol		1–65 μg/dL	
Ferritin	1	2–1,100 ng/mL	
Folate	I	1.3-20.0 ng/mL	
Human chorionic gonadotropin (hCG)	•	5-14,000 mIU/mL	
Vitamin B ₁₂		100-2,200 pg/mL	

^{*}The LN5S CVL will allow Siemens ADVIA (Centaur, XP, and CP) and Atellica IM users to report other major instruments for analytes other than CEA, if needed.

View your expedited linearity evaluations within two business days by logging into e-LAB solutions Suite.

Program Information

- LN5 Eight 4.0-mL liquid serum specimens; appropriate for most major instruments except Siemens ADVIA (Centaur, XP, and CP) and Atellica IM users
- LN5S Thirteen 4.0-mL liquid serum specimens; appropriate for Siemens ADVIA (Centaur, XP, and CP) and Atellica IM users
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Instrumentation Verification Tools

Urine Chemistry Calibration Verification/Linearity LN6			
Analyte	Program Code		
	LN6	LN6 Target Ranges	
Amylase	I	40-2,500 U/L	
Calcium		5-30 mg/dL	
Chloride	ı	20-300 mmol/L	
Creatinine	1	20-540 mg/dL	
Glucose	ı	25-640 mg/dL	
Osmolality	I	30 –1,800 m0sm/kg H ₂ 0	
Phosphorus	I	15–225 mg/dL	
Potassium	I	7–225 mmol/L	
Protein, total	I	10-210 mg/dL	
Sodium	I	20-310 mmol/L	
Urea nitrogen/Urea	I	20-2,000 mg/dL	
Uric acid	I	6-200 mg/dL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Twenty 4.0-mL liquid simulated urine specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Immunology Calibration Verification/Linearity LN7		
Analyte	Program Code	
	LN7	LN7 Target Ranges
Alpha-1 antitrypsin		35-500 mg/dL
Complement C3	ı	21–420 mg/dL
Complement C4	ı	5–125 mg/dL
IgA	ı	32-650 mg/dL
IgG	ı	160-3,800 mg/dL
IgM	ı	25-550 mg/dL
Transferrin	I	50-750 mg/dL

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 2.0-mL liquid serum specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Reproductive Endocrinology Calibration Verification/Linearity LN8		
Analyte	Program Code	
	LN8	LN8 Target Ranges
Estradiol	I	25-4,500 pg/mL
Follicle-stimulating hormone (FSH)	I	3-190 mIU/mL
Human chorionic gonadotropin (hCG)	I	5-8,000 mIU/mL
Luteinizing hormone (LH)	I	2-190 mIU/mL
Progesterone	I	1–50 ng/mL
Prolactin	I	3-315 ng/mL
Testosterone	I	20-1,500 ng/dL

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 4.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Hematology Calibration Verification/Linearity LN9		
Analyte	Program Code	
	LN9	LN9 Target Ranges
Hemoglobin	I	1.0-22.5 g/dL
Platelet count	I	10-4,200 x 10 ⁹ /L
RBC count	I	0.3-7.5 x 10 ¹² /L
WBC count	I	0.5-350.0 x 10 ⁹ /L

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Serum Ethanol Calibration Verification/Linearity LN11			
Analyte Program Code			
LN11 LN11 Target Range			
Serum ethanol	I	15-550 mg/dL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Twenty 3.0-mL liquid specimens
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Program Information

- Seven 3.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

C-reactive Protein Calibration Verification/Linearity LN12			
Analyte Program Code			
LN12 LN12 Target Range			
C-reactive protein	I	7–316 mg/L	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Not appropriate for reporting high-sensitivity C-reactive protein (hsCRP). For reporting hsCRP, use LN21 on page 130.

Program Information

- Seven 1.0-mL liquid serum specimens
- · Two shipments per year

Blood Gas/Critical Care		
Calibration Verification/Linearity	LN13, LN13C	

Ganbration	70111104	cion, Emouri	.,	, =11100
Analyte	Program Code		Program Code	
	LN13	LN13 Target Ranges	LN13C	LN13C Target Ranges
pCO ₂		12-91 mm Hg	1	12-91 mm Hg
рН		6.83-7.82	ı	6.83-7.82
pO ₂	I	18-490 mm Hg	ı	18-490 mm Hg
Calcium, ionized			ı	0.15-3.30 mmol/L
Chloride			ı	62–148 mmol/L
Glucose			ı	10-465 mg/dL
Lactate			ı	0.2-18.0 mmol/L
Potassium			ı	0.5-10.7 mmol/L
Sodium			ı	83–172 mmol/L

Program Information

- LN13, LN13C Ten 2.5-mL ampules of aqueous specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Hemoglobin A1c Accuracy Calibration Verification/Linearity LN15

Analyte	Program Code	
	LN15	LN15 Target Range
Hemoglobin A1c	I	5%-12%

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

CAP-assigned target values are derived from Hemoglobin A1c measurements assayed by National Glycohemoglobin Standardization Program (NGSP) secondary reference laboratories.

Program Information

- Six 0.8-mL liquid human whole blood specimens
- · Two shipments per year

Homocysteine Calibration Verification/Linearity LN16 Analyte Program Code LN16 LN16 Target Range Homocysteine ■ 5-65 μmol/L

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL liquid serum specimens
- · Two shipments per year

Whole Blood Glucose Calibration Verification/Linearity LN17				
Analyte Program Code				
LN17 LN17 Target Range				
Whole blood glucose	I	50-400 mg/dL		

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Five 2.0-mL liquid whole blood specimens
- Report up to 10 different ancillary testing sites or instruments
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Reticulocyte Calibration Verification/Linearity LN19		
Instrument/Method Program Code		
	LN19	LN19 Target Range
Beckman Coulter Unicel DxH series (except DxH 500)	ı	0.5%-28.0%

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Five 2.5-mL liquid whole blood specimens with pierceable caps
- Two shipments per year

Urine Albumin Calibration Verification/Linearity LN20			
Analyte	Program Code		
	LN20	LN20 Target Ranges	
Urine albumin	I	10-350 mg/L	
Urine creatinine	I	20-500 mg/dL	
Urine albumin/creatinine ratio	1		

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

The urine albumin/creatinine ratio results will be evaluated with a calculation verification comparison.

Program Information

- Six 5.0-mL urine specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

CD3+/CD8+

CD3+/CD8+ T lymphocytes absolute

High-Sensitivity C-reactive Protein Calibration Verification/Linearity LN21 Analyte Program Code LN21 LN21 Target Range High-sensitivity C-reactive protein ■ 0.5-18.0 mg/L

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL liquid serum specimens
- For high-sensitivity methods only
- · Two shipments per year

Flow Cytometry Calibration Verification/Linearity LN22			
Analyte	Program Code		
	LN22	LN22 Target Ranges	
CD3+	1	50%-70% positive	
CD3+ T lymphocytes absolute		350–4,000 cells/μL	
CD3+/CD4+		1%-40% positive	
CD3+/CD4+ T lymphocytes absolute	•	6-2.000 cells/uL	

25%-40% positive

250-1,600 cells/μL

Program Information

- Seven 1.0-mL liquid whole blood specimens
- · Two shipments per year

Prostate-Specific Antigen Calibration Verification/Linearity LN23			
Analyte	Program Code		
	LN23	LN23 Target Range	
Prostate-specific antigen	ı	0.1-90.0 ng/mL	

Program Information

- Twelve 1.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Creatinine Accuracy Calibration Verification/Linearity LN24			
Analyte	Program Code		
	LN24	LN24 Target Range	
Creatinine	I	0.6-4.0 mg/dL	
Estimated glomerular filtration rate (eGFR)			

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

eGFR results will be evaluated with a calculation verification comparison.

The College of American Pathologists (CAP) and the National Kidney Disease Education Program (NKDEP) have an initiative to harmonize clinically reported creatinine values. This initiative is analogous to what the federal health agencies and the clinical laboratory community did to improve the accuracy of cholesterol and glycohemoglobin testing.

Troponin I Calibration Verification/Linearity LN25			
Analyte Program Code			
LN25 LN25 Target Range			
Troponin I		0.1-65.0 ng/mL	

LN25 is not appropriate for reporting high-sensitivity troponin. For reporting high-sensitivity troponin I, use program LN48 on page 135.

B-type Natriuretic Peptides Calibration Verification/Linearity LN30		
Analyte	Program Code	
	LN30	LN30 Target Ranges
BNP		18-5,000 pg/mL
NT-proBNP		35-25,000 pg/mL

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL human serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Program Information

- LN25 Seven 2.0-mL liquid serum specimens
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Program Information

- Six 1.0-mL liquid plasma specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Instrumentation Verification Tools

Immunosuppressive Drugs Calibration Verification/Linearity LN31		
Analyte	Program Code	
	LN31	LN31 Target Ranges
Cyclosporine	I	60-1,200 ng/mL
Tacrolimus	I	1.5-30.0 ng/mL

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 2.0-mL liquid whole blood hemolysate specimens
- · Two shipments per year

Ammonia Calibration Verification/Linearity LN32					
Analyte Program Code					
	LN32 LN32 Target Range				
Ammonia		13-900 μmol/L			

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- · Seven 2.0-mL aqueous specimens
- Two shipments per year

Serum Myoglobin Calibration Verification/Linearity LN33			
Analyte Program Code			
	LN33	LN33 Target Range	
Myoglobin		25-900 ng/mL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 1.0-mL liquid serum specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Tumor Markers Calibration Verification/Linearity LN34			
Analyte	Program Code		
	LN34	LN34 Target Ranges	
CA 125	1	1–1,000 U/mL	
CA 15-3	I	2-190 U/mL	
CA 19-9	I	10-900 U/mL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 3.0-mL liquid serum specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Coagulation Calibration Verification/Linearity LN35, LN36, LN37, LN51					
Analyte		Progra	m Code		
	LN35	LN36	LN37	LN51	Target Ranges
Antithrombin activity					10%-130%
Protein C activity					10%-100%
Heparin, low molecular weight					0.1-2.0 U/mL
Heparin, unfractionated					0.1-1.3 U/mL
von Willebrand factor antigen					5%-140%
Factor VIII clot-based				ı	1-200 IU/dL
Factor VIII chromogenic				I	1-200 IU/dL

View your expedited linearity evaluations for LN35, LN36, and LN37 within two business days by logging into e-LAB Solutions Suite.

The LN35, LN36, and LN37 CVL programs meet the CAP Accreditation Checklist requirements HEM.37363, 37365, 37373, and 37375.

Program Information

- LN35, LN37, LN51 Six 1.0-mL frozen plasma specimens per mailing
- LN36 Twelve 1.0-mL frozen plasma specimens per mailing, which include six for low molecular weight heparin and six for unfractionated heparin
- Two shipments per year; ships on dry ice

Viral Load Calibration Verification/Linearity LN38, LN39, LN45, LN52					
Analyte		Progra	m Code		
	LN38	LN39	LN45	LN52	Target Ranges
CMV viral load	ı				316.0-8.0M IU/mL
HIV viral load		ı			50.0-5.0M IU/mL
HCV viral load					50.0-280.0M IU/mL
HBV viral load					1.3 log-8.5 log IU/mL

View your expedited linearity evaluations for LN38, LN39, and LN45 within two business days by logging into e-LAB Solutions Suite.

Program Information

- LN38 Six 1.5-mL liquid plasma specimens
- LN39 Six 2.5-mL liquid plasma specimens
- LN45 Seven 2.5-mL frozen DNA specimens
- LN52 Seven 2.5-mL frozen DNA specimens
- Two shipments per year; LN45 and LN52 ship on dry ice

Instrumentation Verification Tools

11

Vitamin D Calibration Verification/Linearity LN40			
Analyte Program Code			
	LN40	LN40 Target Range	
25-OH vitamin D, total	I	10-135 ng/mL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Procalcitonin Calibration Verification/Linearity LN41			
Analyte Program Code			
	LN41	LN41 Target Range	
Procalcitonin	ı	0.3-175.0 ng/mL	

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL frozen serum specimens
- · Two shipments per year; ships on dry ice

D-dimer Calibration Verification/Linearity LN42						
Analyte	Program Code					
LN42 LN42 Target Range						
D-dimer	I	220-5,500 ng/mL FEU				

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL plasma specimens
- · Two shipments per year

Fibrinogen Calibration Verification/Linearity LN44					
Analyte	Program Code				
LN44 LN44 Target Range					
Fibrinogen	I	80-900 mg/dL			

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL frozen plasma specimens
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year; ships on dry ice

C-peptide/Insulin Calibration Verification/Linearity LN46					
Analyte Program Code					
LN46 LN46 Target Ranges					
C-peptide		0.2-35.0 ng/mL			
Insulin		0.6-800.0 μIU/mL			

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Seven 2.0-mL frozen serum specimens
- Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

High-Sensitivity Troponin T Calibration Verification/Linearity LN47					
Analyte Program Code					
LN47 LN47 Target Range					
High-sensitivity troponin T	ı	10-9,000 ng/L			

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 2.0-mL serum specimens
- Two shipments per year

High-Sensitivity Troponin I Calibration Verification/Linearity LN48				
Analyte	Program Code			
	LN48	LN48 Target Range		
High-sensitivity troponin I		10-25,000 ng/L		

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 2.0-mL serum specimens
- Two shipments per year

Cystatin C Calibration Verification/Linearity LN49			
Analyte/Procedure Program Code			
	LN49	Target Range	
Cystatin C	I	0.5 - 8.0 mg/L	
Estimated glomerular filtration rate (eGFR)	•		

eGFR results will be evaluated with a calculation verification comparison.

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL liquid serum specimens
- · Two shipments per year

Thyroid Panel Calibration Verification/Linearity Lf		N50
Analyte	Program Code	LN50 Target Ranges
	LN50	
Triiodothyronine (T3), free	•	1.0-18.0 pg/mL
Triiodothyronine (T3), total	•	0.4-7.0 ng/mL
Thyroxine (T4), free		0.7–7.0 ng/dL
Thyroxine (T4), total	1	1.0−27.0 µg/dL
Thryoid-stimulating hormone (TSH)	ı	0.1–120.0 μIU/mL

- Eighteen 2.0-mL serum specimens
- Two shipments per year

The CAP is your trusted calibration verification and linearity partner, offering a comprehensive menu of programs for diagnostic confidence.

- Expedited results—View your linearity evaluation for most CVL programs within two business days of data submission.
- Customized report package—Let our team of biostatisticians perform the statistical analysis of your results so you don't have to.
- Objective Assessment Maximize confidence in instrument calibration by using peer group data for a view beyond your laboratory.

See the Instrumentation Verification Tools section of this catalog to determine programs that best fit your laboratory's CVL needs.

Instrumentation Quality Management Programs

Instrumentation I		
Challenges	Program Code	
	I	
Gravimetric pipette calibration	I	
Microtiter plate linearity	I	
Refractometer calibration	I	
Spectrophotometer (stray light check)	I	
Fluorescent intensity check – fluorescent microscopes	I	
pH meter check	I	

WARNING: The Instrumentation (I) program specimens may contain corrosive or toxic substances, environmental hazards, or irritants.

Program	Information

- · Designed to assess instruments not routinely challenged during the proficiency testing process
- · Includes appropriate materials to assess important functional parameters, including accuracy and linearity
- · One shipment per year

Serum Carryover SCO		
Analyte	Program Code	
	SCO	
Creatinine	I	
hCG	I	
Lactate dehydrogenase (LD)	I	
Phenytoin	I	

Program Code
SCO
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ı

Urine Toxicology Carryover UTCO		
Analyte	Program Code	
	UTCO	
Benzoylecgonine	I	
Delta-9-THC-COOH	I	
Opiates	I	
Amphetamine	I	

Program Information

- One 10.0-mL liquid serum specimen (low level) and one 5.0-mL liquid serum specimen (high level)
- · Designed to screen for instrument sample probe carryover
- · One shipment per year

Program Information

- Two 40.0-mL urine specimens (low and high levels)
- Designed to screen for instrument sample probe carryover
- · One shipment per year

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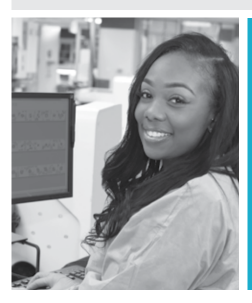
Interfering Substance IFS			
Analyte	Program Code		
	IFS		
	Bilirubin Interferent	Hemoglobin Interferent	Lipid Interferent
Alanine aminotransferase (ALT/SGPT)	ı		ı
Albumin	ı	•	ı
Alkaline phosphatase	I	•	ı
Amylase	I	•	ı
Aspartate aminotransferase (AST/SGOT)	ı		ı
Calcium	I	•	I
Chloride	I		ı
CK-2 (MB) mass	ı	•	I
Creatine kinase (CK)	ı	•	I
Creatinine	ı	•	I
Gamma glutamyl transferase (GGT)	ı	•	I
Glucose	ı	•	I
Iron	I		ı
Lactate dehydrogenase (LD)	ı	•	I
Lipase	ı	•	I
Magnesium	ı	•	I
Osmolality	ı	•	I
Phosphorus	I	•	I
Potassium	ı	ı	ı
Protein, total	I	I	I
Sodium	ı	I	I
Urea nitrogen (BUN)	I	I	I
Uric acid	I	I	I

The material expires December 1, 2025.

Program Information

- Eighteen 10.0-mL liquid serum specimens
- Designed for verifiying manufacturing interference specifications and investigating discrepant results caused by interfering substances
- Submit results any time prior to the material's expiration date.
- One shipment per year

12 Hematology and Clinical Microscopy



When you transmit quantitative PT results directly to the CAP, less equals more.

- Reduce clerical errors and make the PT process more like patient testing.
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Hematology and Clinical Microscopy

Hematology	140
Clinical Microscopy	152

Hematology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Hematology—Basic HE			
Analyte/Procedure	Program Code	Challenges per Shipment	
	HE		
Hematocrit		5	
Hemoglobin		5	
MCV, MCH, and MCHC	I	5	
MPV		5	
Platelet count		5	
RDW		5	
Red blood cell count		5	
White blood cell count		5	

Program Information

- Five 3.0-mL whole blood specimens
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



Hematology Automated Differential Series FH1-FH4, FH9-FH10, FH13, FH16-FH17

Analyte/Procedure	Program Code	Challenges per Shipment
	FH1-FH4, FH9-FH10, FH13, FH16-FH17	
Hematocrit	I	5
Hemoglobin	I	5
Immature granulocyte (IG)	I	5 (FH9 and FH17)
Immature platelet fraction (IPF)/reticulated platelet (RP)	ı	5 (FH9 only)
Large unstained cell (LUC)	I	5 (FH4 only)
MCV, MCH, and MCHC	I	5
MPV	I	5
Nucleated red blood cell count (nRBC)	ı	5 (FH3, FH9, FH13, FH16, and FH17)
Platelet count	I	5
RDW	I	5
Red blood cell count	I	5
White blood cell count	I	5
WBC differential	I	5

For multiple instrument reporting options, see the Quality Cross Check programs, FH3Q, FH4Q, FH9Q, and FH13Q, on page 142.

Program Information

- FH1-4, FH10, FH16-17

 Five 2.5-mL whole blood specimens in vials with pierceable caps
- FH9, FH13 Five 2.0-mL whole blood specimens in vials with pierceable caps
- For method compatibility, see instrument matrix on page 141.
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



Hematology Automated Differential Series, Instrument Matrix

Instrument	FH and FHQ Series								
	FH1	FH2	FH3/ FH3Q	FH4/ FH4Q	FH9/ FH9Q	FH10	FH13/ FH13Q	FH16/ FH3Q	FH17
Abbott Cell-Dyn® 1200, 1400, 1600, 1700, Emerald™									
Horiba ABX 9000+, 9018+, 9020+									
Sysmex K-series, K-1000/KCP-1, KX-21/21N, pocH-100i, XP-series									
CDS/Medonic M-series									
Beckman Coulter® AcT, diff/diff 2™ MD 2/8/10/16, ONYX™, S880, S-plus V, ST, STKR, T-series		•							
Drew Scientific DC-18, I-1800, DREW3, EXCELL 10/16/18									
Horiba ABX Micros									
Mindray BC-2800, 3000/3200 series									
Siemens ADVIA® 360									
Abbott Cell-Dyn 3000, 3500, 3700, 4000, Emerald 22/AL, Ruby™, Sapphire™			•						
Biosystems HA3/HA5			I						
Drew Scientific EXCELL 22, 2280			I						
HumaCount5D			I						
Nihon Kohden MEK 9100									
Orphee Mythic 18, 22 AL, 22 OT, 60									
Siemens ADVIA 560			ı						
Siemens ADVIA 120, 120 w/SP1, 2120				ı					
Abbott Alinity hq, Sysmex XE-2100, XE-2100C, XE-2100D, XE-2100DC, XE-2100D/L (Blood Center), XE-2100L, XE-5000, XN-series (includes RL App), XN-L series, XR-series, XS-500i, XS-800i, XS-1000i, XS-1000i-AL, XS-1000iC, XT-1800i, XT-2000i, XT-4000i, Zybio EXZ 6000 series					1				
Beckman Coulter AcT 5diff (AL, CP, OV)									
DIRUI BF series									
Horiba ABX Pentra 60, 80, 120, Pentra DF Nexus						ı			
Beckman Coulter LH 750, LH 755, LH 780, LH 785, UniCel DxH series (except DxH 500 series)									
Beckman Coulter DxH 500 series									
Horiba Yumizen H500/550, H1500/2500									
Mindray BC-700, BC-720, BC-760, BC-780, BC-6000, BC-6000Plus, BC-6100, BC-6100Plus, BC-6200, BC-6200Plus, BC-6600, BC-6600Plus, BC-6700, BC-6800, BC-6800Plus, BC-7500 CRP									

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Quality Cross Check—Hematology FH3Q, FH4Q, FH9Q, FH13Q						
Analyte/Procedure		Challenges per Shipment				
	FH3Q	FH4Q	FH9Q	FH13Q		
Hematocrit		•			3	
Hemoglobin		ı			3	
Immature granulocyte (IG)					3	
Immature platelet fraction (IPF)%					3	
Large unstained cells (LUC)		•			3	
MCV, MCH, MCHC		•			3	
MPV		•			3	
Nucleated red blood cell count (nRBC)	ı			I	3	
Platelet count		•			3	
RDW		•			3	
Red blood cell count		•			3	
WBC differential		•			3	
White blood cell count					3	

These programs do not meet regulatory requirements for proficiency testing; see the FH Series on page 140. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- · Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Blood Cell Identification, Photographs BCP					
Procedure	Program Code	Challenges per Shipment			
	ВСР				
Blood cell identification		5			
Educational challenge(s)		5			

Blood Cell Identification, Virtual BCPV Analyte/Procedure **Program Code** Challenges per Shipment **BCPV** ı 5 **Blood cell identification** 5 Educational challenge ı

Program Information

- FH3Q, FH4Q, FH9Q, FH13Q -Three 2.5-mL whole blood specimens in vials with pierceable caps
- · Report up to three instruments.
- · For method compatibility, see instrument matrix on page 141.
- · Conventional and International System of Units (SI) reporting offered
- · Two shipments per year

Program Information

- Ten images, each available as photographs and online images
- · Three shipments per year



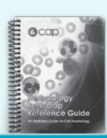
Program Information

- Ten online images
- Three shipments per year



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Assessment of Consistency of Peripheral Blood Morphologic Observations QPC10/QPC25

Introduction

The widespread use of automated white blood cell (WBC) differential counts and computer-generated whole slide imaging has decreased the time that the medical laboratory scientist/technologist staff dedicate to morphological assessment of blood cells. However, these staff must maintain their morphological skills. Laboratories have an annual requirement to do a morphologic comparison of their technical staff's peripheral blood smear results, assess their competency on peripheral blood smears, and provide appropriate education.

Objectives

This study will help assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the performance of accurate WBC differential counts and other peripheral blood smear morphological assessments. The evaluation provided will assist in the construction of individual educational programs for the technical staff and show areas that need focused review and improvement. The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for consistency of reporting morphology among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

A series of online, whole slide images of Wright or Wright-Giemsa stained peripheral blood smears using DigitalScope® technology will be available to each participating institution to assess technologists' performance on WBC differential counts and morphology assessment. Each technologist will receive their own kit. Technologists will provide information about their continuing education and professional background. Information will be collected from each site regarding their institution's minimum continuing education requirements for their technologists in hematology and relevant procedures and policies related to peripheral blood smear assessment.

Performance Indicators

- Individual technologist score (%) based on a standardized competency assessment method to determine a technologist's ability to identify various WBC types, red blood cell morphology, and platelet morphology in normal and abnormal cases
- · Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your staff technical competency assessment requirements:

- Result forms for up to 10 technologists (QPC10)
- Result forms for up to 25 technologists (QPC25)
- Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

Preliminary study reports are provided at institution and technologist levels.

*Participation in this study helps laboratories meet applicable requirements:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Checklist statements: HEM.34400, consistency of morphologic observation among personnel performing blood cell microscopy at least annually; GEN.55500, element 5, Competency Assessment of Testing Personnel; GEN.55525, Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed by a qualified individual
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, Pl.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members.

This is a one-time study conducted in the third quarter.

Blood Parasite BP				
Procedure	Program Code	Challenges per Shipment		
	ВР			
Blood parasite identification (thin/thick film sets*)	ı	5		

^{*}This program will include corresponding thick films when available.

- Five Giemsa-stained blood film sets, photographs, and/or online images
- Percent parasitemia reporting is provided when appropriate for educational purposes.
- A variety of blood parasites, including Plasmodium, Babesia, Trypanosoma, and filarial worms
- Three shipments per year

Bone Marrow Cell Differential BMD				
Procedure	Program Code	Challenges per Shipment		
	BMD			
Bone marrow differential		1		
Bone marrow cell identification		5		

Additional Information

- Examine an online whole slide image that includes a manual 500-count bone marrow differential and annotated cells for identification.
- Recognize and integrate problem-solving skills through the use of interpretive questions found throughout the discussion.
- Evaluate cell morphology and identify specific cells in bone marrow.
- See system requirements on page 12.

Erythrocyte Sedimentation Rate ESR, ESR1, ESR2, ESR3					
Procedure Program Code Challenges per Shipm					Challenges per Shipment
	ESR	ESR1	ESR2	ESR3	
All methods except the ALCOR, Alifax®, Sedimat 15®, and Sedimat 15 Plus					3
Sedimat 15, Sedimat 15 Plus		ı			3
Alifax			•		3
Mindray BC 700 series, Mindray BC 6800 Plus, Mindray BC 7600/7800/7900, and Mindray BP 200n series					3
ALCOR iSED®, miniiSED®				I	3

Program Information

- One online bone marrow aspirate whole slide image that includes five annotated cells for identification
- Powered by DigitalScope® technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



- ESR, ESR1 Three 6.0-mL whole blood specimens
- ESR2 Three 3.0-mL latex bead specimens
- ESR3 Three 3.5-mL whole blood specimens
- Two shipments per year

Fetal Red Cell Detection HBF					
Procedure Program Code Challenges per Shipm					
	HBF				
Kleihauer-Betke and flow cytometry	1	2			
Rosette fetal screen	I	2			
Acid elution whole slide image	ı	1			

- Two 1.2-mL liquid whole blood specimens
- Not designed for F-cell quantitation
- Two online whole slide images per year with optional grids for cell counting
- Powered by DigitalScope technology
- Two shipments per year

Reticulocyte Series RT, RT2, RT3, RT4					
Instrument/Method	Program Code			Challenges per Shipment	
	RT	RT2	RT3	RT4	
Abbott Alinity hq, Abbott Cell-Dyn 4000, Sapphire, Siemens ADVIA 120/2120, and all other automated and manual methods	ı				3
Abbott Cell-Dyn 3500, 3700, Ruby		ı			3
Beckman Coulter LH 500, LH 700 series, UniCel DxH series					3
Sysmex XE-2100, XE-2100C, XE-2100D, XE-2100DC, XE-2100L, XE-5000, XN-L series, XN-series (includes RL App), XR-series, XT-2000i, XT-4000i				•	3
Pierceable caps				I	3

For specific program testing components, see reticulocyte matrix below.

Program Information

- RT, RT2 Three 1.0-mL stabilized red blood cell specimens
- RT3, RT4 Three 3.0-mL stabilized red blood cell specimens
- Two shipments per year

Reticulocyte, Matrix

Program Code	Reticulocyte count, percent	Absolute reticulocyte count	Immature Reticulocyte Fraction (IRF)	Reticulocyte Hemoglobin Concentration (CHr)	Reticulocyte Hemoglobin (RET-He)
RT/RTQ			ı		
RT2					
RT3/RT3Q			ı		
RT4/RT4Q	I		I		1

Quality Cross Check—Reticulocyte RTQ, RT3Q, RT4Q				
Instrument/Method	strument/Method Program Code			Challenges per Shipment
	RTQ	RT3Q	RT4Q	
Abbott Alinity hq, Abbott Cell-Dyn 4000, Sapphire, Siemens ADVIA 120/2120, and all other automated and manual methods	•			3
Beckman Coulter, LH 500, LH 700 series, UniCel DxH series				3
Sysmex XE-2100, XE-2100C, XE-2100D, XE-2100DC, XE-2100L, XE-5000, XN-L series, XN-series (includes RL App), XR-series, XT-2000i, XT-4000i			ı	3

These programs do not meet regulatory requirements for proficiency testing; see the RT Series on page 146. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

Hemoglobinopathy HG				
Procedure	Program Code	Challenges per Shipment		
	HG			
Hemoglobin identification and quantification	ı	4		
Educational dry challenges	1	2		
Hemoglobin A2 quantitation	1	4		
Hemoglobin F quantitation	1	1		
Sickling test, qualitative	1	4		

Rapid Total White Blood Cell Count RWBC					
Procedure Program Code Challenges per Shipment					
RWBC					
Rapid total white blood cell count	•	5			

Program Information

- RTQ Three 1.0-mL stabilized red blood cell specimens
- RT3Q, RT4Q Three 3.0-mL stabilized red blood cell specimens
- Includes percentage and absolute result reporting
- Report up to three instruments.
- Two shipments per year

Program Information

- Four 0.5-mL stabilized red blood cell specimens
- Two educational dry challenges (case histories, electrophoresis patterns, and clinical interpretation questions)
- Two shipments per year

- Five 2.0-mL whole blood specimens
- For use with the HemoCue WBC instrument
- Three shipments per year

Sickle Cell Screening SCS					
Procedure Program Code Challenges per Shipme					
scs					
1	3				
	Program Code				

Transfusion-Related Cell Count TRC					
Procedure Program Code Challenges per Shipme					
	TRC				
Platelet count (platelet-rich plasma)	ı	5			
WBC count	I	4			
Dry challenge		2			

WBC counts must be performed using a Nageotte chamber, by fluorescence microscopy, or by flow cytometry.

Waived Combination HCC					
Analyte Program Code Challenges per Shipmer					
HCC					
Hemoglobin		2			
Whole blood glucose		2			

Waived Hemog	NEW	
Analyte	Program Code	Challenges per Shipment
	HCC1	
Hemoglobin	1	2

Program Information

- Three 1.0-mL whole blood specimens
- Two shipments per year

Program Information

- Five 1.2-mL suspensions of platelet-rich plasma
- Two 1.0-mL vials leukocytereduced platelet material
- Two 1.0-mL vials leukocytereduced red blood cells
- Three shipments per year

Program Information

- Two 1.5-mL whole blood specimens; two shipments per year
- · Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix on page 65.

- Two 1.0-mL whole blood specimens; two shipments per year
- · Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix on page 65.

Waived Combination HCC2				
Analyte Program Code Challenges per				
	HCC2			
Hematocrit		2		
Hemoglobin	2			
Urinalysis/urine hCG		2		
Whole blood glucose		3		

- Total of four shipments per year
- Hematocrit, hemoglobin, and urinalysis/urine hCG testing
 Two 3.0-mL whole blood specimens and two 10.0-mL urine specimens; two shipments per year: A and C
- Whole blood glucose testing

 Three 2.0-mL whole blood specimens; two shipments per year: B and D
- Conventional and International System of Units (SI) reporting offered
- To identify instrument specific programs, refer to the whole blood chemistry compatibility matrix on page 65.

Virtual Peripheral Blood Smear VPBS				
Procedure	Challenges per Shipment			
	VPBS			
WBC differential	I	3		
Platelet estimate	I	3		
RBC morphology	I	3		
Blood cell identification	1	15		

Additional Information

- Examine online whole slide images that include a manual 100 white blood cell (WBC) differential count and annotated cells for identification.
- Evaluate and identify red blood cell (RBC) morphology and identify specific white blood cells (WBC) in peripheral blood.
- Recognize and integrate problem-solving skills through the use of interpretive questions found throughout the discussion.
- · See system requirements on page 12.

- Three online peripheral blood whole slide images that include 15 annotated cells for identification
- Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Expanded Virtual Peripheral Blood Smear EHE1					
Procedure Program Code Challenges per Shipme					
	EHE1				
WBC differential	I	2			
Platelet estimate	I	2			
RBC morphology	I	2			
Blood cell identification	ı	10			

Additional Information

- More challenging and/or complex testing than the Virtual Peripheral Blood Smear (VPBS) program
- Examine online whole slide images that include a manual 100 white blood cell (WBC) differential count and annotated cells for identification.
- · Comprehensive case studies
- Recognize and integrate problem-solving skills through the use of interpretive questions found throughout the discussion.
- Evaluate and identify red blood cell (RBC) morphology and identify specific white blood cells (WBC) in peripheral blood.
- See system requirements on page 12.

- Two online peripheral blood whole slide images that include 10 annotated cells for identification
- Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Hematopathology Online Education HPATH/HPATH1					
Program Code Challenges per Shipment					
HPATH/HPATH1					
Hematopathology online case review					

Additional Information

HPATH/HPATH1 prepares pathologists and pathology trainees with an interest in hematopathology to succeed by providing ongoing diagnostic learning and self-assessment.

The program includes 10 challenge cases per year written by expert hematopathologists. For each case the learner will:

- Review virtual whole slide image(s) in the context of a clinical scenario and relevant laboratory data.
- Select and interpret appropriate ancillary studies (IHC, flow cytometry, FISH, cytogenetics, molecular studies, etc) with real-time feedback.
- Simulate a real-world diagnostic workup to arrive at the appropriate diagnosis.
- · Apply knowledge to answer three CME questions.

Cases span the breadth of benign and neoplastic hematopathology. Discussions incorporate current best practices in the workup, diagnosis, and subclassification of hematolymphoid diseases.

Neoplastic entities are discussed in the context of both the International Consensus Classification (ICC) and the fifth edition of the World Health Organization (WHO) classification.

See system requirements on page 12.

Program Information

- HPATH Five diagnostic challenges per activity (two activities per year), with online whole slide images. Reporting with CME credit is available for one participant.
- HPATH1 Reporting option with CME credit for each additional participant (within the same institution); must order in conjunction with program HPATH
- Earn a maximum of 12.5 CME credits (AMA PRA Category 1 Credits™) per participant.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available

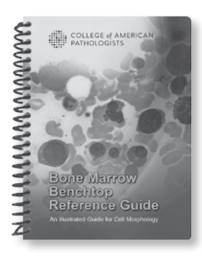


Bone Marrow Benchtop Reference Guide

With more than 60 different identifications and a detailed description for each cell morphology, this illustrated guide is an affordable, convenient way to identify various cell types quickly and confidently. Plus, its rugged construction makes it well-suited for heavy use at the benchtop.

Add it to your order, or view sample pages and purchase online.

- printed books at estore.cap.org
- ebooks at ebooks.cap.org



Item number: BMBRG

Spiral bound; 66 pages; 2019

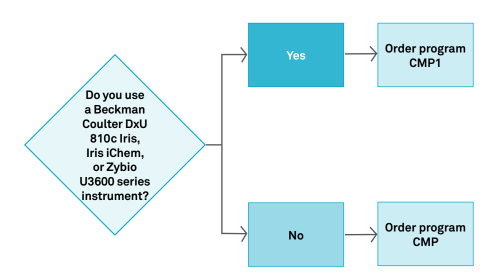
Clinical Microscopy

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Urinalysis and Clinical Microscopy CMP, CMP1				
Analyte/Procedure	Progra	m Code	Challenges per Shipment	
	СМР	CMP1		
Bilirubin			3	
Blood or hemoglobin			3	
Body fluid photographs		•	3	
Glucose		•	3	
hCG urine, qualitative		•	3	
Ketones	•	1	3	
Leukocyte esterase	•	•	3	
Nitrite	•	1	3	
Osmolality		I	3	
рН	•	•	3	
Protein, qualitative	•	1	3	
Reducing substances	•	1	3	
Specific gravity		•	3	
Urine sediment photographs	•	ı	3	
Urobilinogen	I		3	

For multiple instrument reporting options, see the Quality Cross Check program, CMQ, on page 153.

- CMP Three 10.0-mL liquid urine specimens; for use with all instruments except Beckman Coulter DxU 810c Iris and Iris iChem; six images, each available as photographs and online images
- CMP1 Three 10.0-mL liquid urine specimens; for use with Beckman Coulter DxU 810c Iris, Iris iChem, and Zybio U3600 Series instruments only, urinalysis; six images, each available as photographs and online images
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year



Quality Cross Check—Urinalysis CMQ				
Analyte	Program Code Challenges per Shi			
	СМQ			
Bilirubin	ı	3		
Blood or hemoglobin	ı	3		
Glucose	I	3		
hCG urine, qualitative	ı	3		
Ketones	I	3		
Leukocyte esterase	ı	3		
Nitrite	ı	3		
Osmolality	ı	3		
рН	ı	3		
Protein, qualitative	ı	3		
Reducing substances	ı	3		
Specific gravity	ı	3		
Urobilinogen	I	3		

This program does not meet regulatory requirements for proficiency testing; see programs CMP and CMP1 on page 152. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments, and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

Clinical Microscopy Miscellaneous Photopage CMMP				
Procedure	Program Code Challenges per Shi			
	СММР			
Fern test (vaginal)	I	1		
KOH preparation (skin)	I	1		
Nasal smear	I	1		
Pinworm preparation	I	1		
Spermatozoa	I	1		
Stool for leukocytes	I	1		
Urine sediment photographs	I	3		
Vaginal wet preparation photographs (for clue cells, epithelial cells, trichomonas, or yeast)	ı	1		

Program Information

- Three 10.0-mL liquid urine specimens for use with all instruments
- Report up to three instruments.
- Two shipments per year

- Ten images, each available as photographs and online images
- Two shipments per year

Amniotic Fluid Leakage AFL					
Procedure	Program Code	Challenges per Shipment			
AFL					
pH interpretation	3				

- · Three 2.0-mL liquid specimens
- For use with nitrazine paper and the Amniotest™
- Two shipments per year

Automated Body Fluid Series ABF1, ABF2, ABF3				
Procedure	Program Code Challenges per Shipment			
	ABF1	ABF2	ABF3	
Red blood cell fluid count				2
Total nucleated cell/WBC fluid count				2

For method compatibility, see automated body fluid instrument matrix below.

Program Information

- ABF1-3 Two 3.0-mL simulated body fluid specimens
- · Two shipments per year

Automated Body Fluid, Instrument Matrix

Instrument	ABF Series		
	ABF1	ABF2	ABF3
Advanced Instruments GloCyte, Siemens ADVIA 120/2120 series			
Beckman Coulter LH 700 series, Unicel DxH series			
Sysmex XE-2100, XE-2100C, XE-2100D, XE-2100DC, XE-2100L, XE-5000, XN-series, XN-L series, XT-1800i, XT-2000i, XT-4000i			
Beckman Coulter iQ200/DxU Iris series			•

Virtual Body Fluid VBF					
Procedure Program Code Challenges per Shipmen					
VBF					
Body fluid cell differential	I	2			
Body fluid cell identification	I	10			

Additional Information

- Examine online whole slide images that include a manual differential count and annotated cells for identification.
- Evaluate cell morphology and identify specific cells in a body fluid.
- See system requirements on page 12.

- Two online whole slide body fluid images that include 10 annotated cells for identification
- · Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Assessment of Consistency of Body Fluid Morphologic Observations QPB10/QPB25

Introduction

Laboratories receive a variety of body fluids for evaluation that technologists review. Technical staff must maintain their identification skills of these specimens, and laboratories are required to provide education and to assess consistency of reporting morphology among staff and competency of body fluid cell identification on an annual basis.

Objectives

This study will assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the performance of accurate body fluid cell counts and identification of other body fluid features. Results of this study will assist individuals, the laboratory director, and the manager with areas to focus on for improvement and education.

The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for consistency of reporting morphology among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

Technologists will access a series of online, whole slide images to assess their ability to perform cell differentials on Wright-stained body fluids and to identify miscellaneous cells and inclusions in cytocentrifuged preparations using their own kit and result form. Participants will provide additional information about their competency assessment programs, continuing education, and professional background. Information will be collected from each site regarding their institution's minimum continuing education programs and requirements for their technologists in who review body fluids, and relevant procedures and policies related to body fluid review assessment.

Information will be collected from each site regarding their institution's minimum continuing education requirements for their technologists in hematology and relevant procedures and policies related to peripheral blood smear assessment.

Performance Indicators

- Individual technologist score (%) based on a standardized competency assessment method to determine a technologist's ability to identify various white blood cell types, microorganisms, and other cells and inclusions present in normal and abnormal cases in comparison to consensus responses
- Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your technical staff morphology and competency assessment requirements:

- Result forms for up to 10 technologists (QPB10)
- Result forms for up to 25 technologists (QPB25)
- · Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

*Participation in this study helps laboratories meet applicable requirements:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Checklist statements: HEM.35566, consistency of morphologic observation among personnel performing body fluid cell differentials at least annually; GEN.55500, Competency Assessment of Testing Personnel (element 5); GEN.55525, Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed by a qualified individual
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, PI.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members

This is a one-time study conducted in the second quarter.

Automated Urine Microscopy UAA, UAA1			
Analyte	Program Code Challenges per Shipme		Challenges per Shipment
	UAA	UAA1	
Casts, quantitative/qualitative	ı	•	2
Crystals, quantitative/qualitative	ı		2
Epithelial cells, quantitative/ qualitative		ı	2
Red blood cells, quantitative/ qualitative	ı	ı	2
White blood cells, quantitative/ qualitative	ı	I	2

For method compatibility, see automated urine microscopy instrument matrix below.

Do you use a Beckman Coulter iQ200/DxU Iris series microscopic analyzer, a DIRUI FUS-200 system, a Roche cobas u701, or a Zybio U3600 series instrument? No Order program UAA1

Automated Urine Microscopy, Instrument Matrix

Instrument	UAA, UAA1	
	UAA	UAA1
Beckman Coulter iQ200/DxU Iris series		
DIRUI FUS-200	I	
Roche cobas u701	1	
77 Elektronika		I
Zybio U3600 series	I	
ARKRAY Aution Hybrid		
Siemens Atellica UAS 800		I
Sysmex UF 50, 100, 500i, 1000i, 3000/4000/5000, Sysmex UX 2000		ı

- UAA Two 10.0-mL liquid urine specimens for use with Beckman Coulter Iris, DIRUI, Roche, and Zybio instruments
- UAA1 Two 12.0-mL liquid urine specimens for use with Sysmex instruments
- Two shipments per year

Crystals BCR, BFC, URC				
Procedure	Pr	ogram Co	de	Challenges per Shipment
	BCR	BFC	URC	
Bile crystal identification	I			2
Body fluid crystal identification				2
Urine crystal identification			ı	2

body fluid specimens (eg, synovial fluid)

Program Information • BCR - Two photographs • BFC - Two 1.5-mL simulated

- URC Two 1.5-mL urine specimens
- Two shipments per year

Dipstick Confirmatory DSC				
Analyte Program Code Challenges per Shipme				
DSC				
Bilirubin	I	2		
Protein	I	2		

Program Information

- Two 12.0-mL liquid urine specimens
- For use with methods to confirm positive bilirubin and protein dipstick results
- · Two shipments per year

Fecal Fat FCFS				
Analyte Program Code Challenges per Shipme				
FCFS				
Fecal fat, qualitative ■ 2				

Program Information

- Two 10.0-g simulated fecal fat specimens
- For microscopic detection of neutral fats (triglycerides) and/or split fats (total free fatty acids)
- Two shipments per year

Fetal Hemoglobin APT			
Analyte Program Code Challenges per Shipme			
	APT		
Fetal hemoglobin (gastric fluid or stool)	1	2	

Program Information

- Two 1.2-mL simulated body fluid specimens
- Two shipments per year

Gastric Occult Blood GOCB				
Analyte Program Code Challenges per Shipmer				
GOCB				
Gastric occult blood	I	3		
Gastric pH ■ 3				

- Three 2.0-mL simulated gastric fluid specimens
- Two shipments per year

Glucose-6-Phosphate Dehydrogenase G6PDS			
Analyte Program Code Challenges per Shipment			
G6PDS			
ı	2		

Hemocytome	eter Fluid Count	HFC
Procedure	Program Code	Challenges per Shipment
	HFC	
Cytopreparation differential		3
Red blood cell fluid count		3

This program has limited stability. Laboratories outside the US or Canada should consider purchase of HFCI, which has longer stability.

Program Information

- Two 0.5-mL lyophilized hemolysate specimens
- · Two shipments per year

Program Information

Program Information

5-part differential · Powered by DigitalScope

• Designed for laboratories

that have experienced

· Two shipments per year

significant shipping and

receiving issues and need longer program stability

outside the US or Canada

technology

• Three 2.0-mL simulated body fluid specimens; two online whole slide images for 2- and

- Three 1.0-mL simulated body fluid specimens
- · Two shipments per year

Hemocytometer Fluid Count, International HFCI			
Procedure	Program Code	Challenges per Shipment	
	HFCI		
Body fluid differential	I	2	
Red blood cell fluid count	I	3	
Total nucleated cell/WBC fluid count	I	3	

Additional Information

Total nucleated cell/WBC

fluid count

- This program meets the CAP's Laboratory Accreditation Program requirements.
- Examine online whole slide images that include a manuaøl differential count.
- See system requirements on page 12.

Hemocytometer Fluid Count, International HFCI			
Procedure	Program Code	Challenges per Shipment	
	HFCI		
Body fluid differential	I	2	
Red blood cell fluid count	•	3	
Total nucleated cell/WBC fluid count	ı	3	

- Three 2.0-mL simulated amniotic fluid specimens
- For use with LBC methods performed on all hematology analyzers
- · Two shipments per year

Lamellar Body Count LBC							
Procedure	Program Code Challenges per Shipment						
	LBC						
Lamellar body count	1	3					

Occult Blood OCB					
Analyte	Program Code	Challenges per Shipment			
	ОСВ				
Occult blood	1	3			

For multiple instrument reporting options, see the Quality Cross Check program, OCBQ, below.

Program Information

- Three 2.0-mL simulated fecal specimens
- · Two shipments per year

Quality Cross Check—Occult Blood OCBQ						
Analyte	Program Code Challenges per Shipmet					
	OCBQ					
Occult blood	ı	3				

This program does not meet regulatory requirements for proficiency testing; see program OCB, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Fetal Membranes/Preterm Labor ROM1						
Procedure Program Code Challenges per Shipr						
	ROM1					
Fetal membranes/preterm labor		3				

Special Clinical Microscopy SCM1, SCM2						
Analyte/Procedure	Program Code Challenges per Shipmen					
	SCM1	SCM2				
Urine hemosiderin, Prussian blue			3			
Urine eosinophils, Wright stain			3			

Program Information

- Three 2.0-mL simulated fecal specimens
- Report up to three instruments.
- · Two shipments per year

Program Information

- Three 0.5-mL simulated vaginal specimens for methods such as Actim PROM, AmniSure, Clinical Innovations, and PartoSure
- · Two shipments per year

- SCM1, SCM2 Three images, each available as photographs and online images
- · Two shipments per year

12

Ticks, Mites, and Other Arthropods TMO					
Procedure	Program Code	Challenges per Shipment			
	ТМО				
Tick, mite, and arthropod identification	I	3			

Urine hCG UHCG						
Procedure	Program Code	Challenges per Shipment				
	UHCG					
Urine hCG, qualitative	ı	5				

Program Information

- Three images, each available as photographs and online images
- Two shipments per year

Program Information

- Five 1.0-mL urine specimens
- Three shipments per year

Urine Albumin and Creatinine, Semiquant UMC							
Analyte/Procedure	Program Code	Challenges per Shipment					
	UMC						
Creatinine, semiquantitative	1	2					
Urine albumin (microalbumin): creatinine ratio	ı	2					
Urine albumin (microalbumin), semiquantitative/qualitative	ı	2					

For quantitative reporting, refer to program U, page 68.

Progra	am	Inf	forr	ma	tio	n
_	_	^				

- Two 3.0-mL liquid urine specimens
- For use with dipstick and semiquantitative methods only
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year

Worm Identification WID					
Procedure	Program Code	Challenges per Shipment			
	WID				
Worm identification		3			

- Three images, each available as photographs and online images
- Two shipments per year

13

13 Reproductive Medicine



Enhance your learning with continuing education (CE) content included with many of our proficiency testing programs.

- For many of our PT/EQA programs, each member of a participating laboratory has complimentary enrollment to online CE activities.
- Advance skills with education activities developed by more than 600 physicians and doctoral scientists with expertise in pathology and laboratory medicine.

Reproductive Medicine

Andrology and Embryology......162

Andrology and Embryology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Semen Analysis SC, SC1, PV, PV1, SM, SV, ASA								
Procedure		Program Code					Challenges per Shipment	
	sc	SC1	PV	PV1	SM	sv	ASA	
Sperm count and presence/ absence (manual methods)								2
Sperm count (automated methods)		•						2
Postvasectomy sperm count and presence/absence (manual methods)			•					2
Postvasectomy sperm count (automated methods)				•				2
Sperm morphology								2
Sperm viability						I		2
Antisperm antibody IgG							I	2

Program Information

- SC Two 0.3-mL stabilized sperm specimens
- SC1 Two 1.0-mL stabilized sperm specimens
- PV Two 0.3-mL stabilized sperm specimens with counts appropriate for postvasectomy testing
- PV1 Two 1.0-mL stabilized sperm specimens with counts appropriate for postvasectomy testing
- SM Two prepared slides for staining
- SV Two eosin-nigrosinstained slides
- ASA Two 0.3-mL serum specimens
- · Two shipments per year



Sperm Count, Motility, Morphology, and Viability SMCD, SM1CD, SM2CD

Procedure		Program Code				
	SMCD	SM1CD	SM2CD			
Sperm count	1			2		
Sperm motility/forward progression	I			2		
Sperm classification		ı		10		
Sperm morphology				2		
Sperm viability			•	2		

- · SMCD Online video clips of sperm available for hemocytometer, Makler, and disposable chambers
- SM1CD, SM2CD Two online challenges that may be viewed as whole slide images powered by DigitalScope® technology
- · Two online activites per year; your CAP shipping contact will be notified via email when the activity is available



Embryology EMB						
Procedure Program Code Challenges per						
	ЕМВ					
Embryo transfer and quality assessment (three- and five-day-old embryos)	ı	4				

- Two online sets of five video clips
- Two online activites per year; your CAP shipping contact will be notified via email when the activity is available



Sex Hormones Y/YY						
Analyte	Program Code	Challenges per Shipment				
	Y/YY					
11-deoxycortisol	1	5				
17-hydroxyprogesterone	1	5				
Androstenedione	1	5				
DHEA sulfate		5				
Estradiol	•	5				
Estriol, unconjugated (uE3)	•	5				
Follicle-stimulating hormone (FSH)		5				
Growth hormone (GH)	•	5				
IGF-1 (somatomedin C)	•	5				
Luteinizing hormone (LH)		5				
Progesterone		5				
Prolactin		5				
Sex hormone-binding globulin (SHBG)		5				
Testosterone		5				
Testosterone, bioavailable (measured)		5				
Testosterone, free (measured)		5				

Program Information

- Y Five 5.0-mL liquid serum specimens in duplicate
- YY Five 5.0-mL liquid serum specimens in triplicate
- · Conventional and International System of Units (SI) reporting offered
- Three shipments per year



Antimüllerian Hormone AMH Analyte **Program Code** Challenges per Shipment **AMH** Antimüllerian hormone 3

- Three 1.0-mL lyophilized serum specimens
- Two shipments per year

Performance Analytics Dashboard: Bringing it all together



The CAP's Performance Analytics Dashboard provides valuable insights into your laboratory's performance with a single comprehensive view of your CAP PT results and accreditation status.

Simplify analysis and reporting of PT performance data

- Quickly spot unacceptable results for follow-up to mitigate risk of inaccurate patient test results
- Review three years of PT results to identify trends and early indicators of potential problems

Prepare for your next CAP accreditation inspection

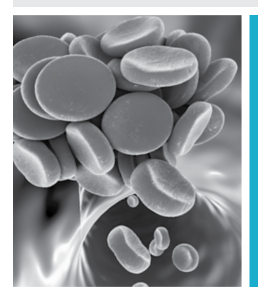
- Manage risk and compliance by identifying areas of improvement based on past deficiencies
- Review PT performance data to ensure appropriate corrective action has been taken for each unacceptable result

Monitor performance of your laboratory or system from a single dashboard

- Benchmark laboratory performance
- Export PT performance from individual laboratories or across the system for quality review meetings

View your laboratory's Performance Analytics Dashboard by accessing e-LAB Solutions Suite (ELSS) from cap.org.

14 Coagulation



Provide for patient care and safety.

The CAP continues to support laboratory quality initiatives through the development, maintenance, and enhancement of effective PT/EQA programs for coagulation.

Coagulation

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Coagulation—Limited CGB, CGL, CGDF						
Analyte	Program Code Challenges per Shipment					
	CGB	CGL				
Activated partial thromboplastin time		•		5		
Fibrinogen		ı		5		
International normalized ratio (INR)*		•		5		
Prothrombin time		•		5		
D-dimer		ı	•	2		
Fibrin(ogen) degradation products, plasma		ı	•	1		
Fibrin(ogen) degradation products, serum		I		1		
Fibrin monomer		I		2		

^{*}Participants reporting INR results will receive a special evaluation to assess the INR calculation. For multiple instrument reporting options, see the Quality Cross Check program, CGLQ, below.

Program Information

- CGB Five 1.0-mL lyophilized plasma specimens
- CGL Seven 1.0-mL lyophilized plasma specimens and one 2.0-mL serum specimen
- CGDF One 2.0-mL serum specimen; two 1.0-mL lyophilized plasma specimens
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



Quality Cross Check—Coagulation CGLQ						
Analyte	Program Code	Challenges per Shipment				
	CGLQ					
Activated partial thromboplastin time	1	3				
Fibrinogen	ı	3				
Prothrombin time	I	3				
D-dimer	I	2				
Fibrin(ogen) degradation products, plasma	ı	1				
Fibrin(ogen) degradation products, serum	I	1				

This program does not meet regulatory requirements for proficiency testing; see program CGL, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

- Three 1.0-mL lyophilized plasma specimens in triplicate, two 1.0-mL lyophilized plasma specimens, and one 2.0-mL serum specimen
- Report up to three instruments.
- · Two shipments per year

Coagulation—Extended CGE/CGEX						
Analyte Program Code Challenges per Shipment						
	CGE/CGEX					
See analyte listing below		2				

- CGE Two 1.0-mL lyophilized plasma specimens (three vials each)
- CGEX Two 1.0-mL lyophilized plasma specimens (five vials each)
- Two shipments per year

Coagulation Analyte Listing (Quantitative Results)

50:50 mixing study, PT and aPTT Plasminogen activator inhibitor
Activated partial thromboplastin time Plasminogen activity/antigen

Activated protein C resistance Prekallikrein
Alpha-2-antiplasmin Protein C
Antithrombin activity/antigen Protein S

Dilute prothrombin time

Factors II, V, VII, VIII, IX, X, XI, XII, and XIII

Fibrinogen antigen

Thrombin time

Heparin-induced thrombocytopenia (HIT)

Expanded Coagulation Factors ECF							
Analyte/Procedure	Program Code	Challenges per Shipment					
	ECF						
Factor II		3					
Factor V		3					
Factor VII		3					
Factor VIII clot based	I	3					
Factor VIII chromogenic		3					
Factor IX		3					
Factor IX chromogenic	I	3					
Factor X clot based		3					
Factor X chromogenic		3					
Factor XI	I	3					
Factor XII		3					
Factor XIII		3					
Fibrinogen antigen		3					
Reptilase time		3					
Thrombin time	I	3					

- Three 1.0-mL lyophilized plasma specimens (three vials each)
- Two shipments per year

Coagulation Special Testing Series CGS1, CGS2, CGS3, CGS4, CGS5, CGS7

CGS3, CGS4, CGS5, CGS7							
Module/Analyte		Challenges per Shipment					
		Program Code					
	CGS1	CGS2	CGS3	CGS4	CGS5	CGS7	
Activated partial thromboplastin time*	2		2	3			
International normalized ratio (INR)	2			3			
Prothrombin time*	2			3			
Lupus Anticoagulant and Mixing St	udies M	odule					
Dilute prothrombin time	2						
Dilute Russell's viper venom time	2						
Lupus anticoagulant sensitive aPTT (confirmation and screen)	2						
50:50 mixing studies, PT and aPTT	2						
Thrombophilia Module	ı			1	I		
Activated protein C resistance		2					
Antithrombin (activity, antigen)		2					
Protein C (activity, antigen)		2					
Protein S (activity, free antigen, total antigen)		2					
von Willebrand Factor Antigen Mod	lule	ı	ı	J			
Factor VIII assay			2				
von Willebrand factor (antigen, activity, multimers)			2				
Factor VIII inhibitor			2				
Heparin Module				ı			
Heparin activities using methodologies including Anti-Xa (unfractionated, low molecular weight, and hybrid curve)				3			
Thrombin time				3			
Heparin-Induced Thrombocytopen	ia Modul	le					
Appropriate with methods such as Immucor Lifecodes PF4 IgG and Immucor Lifecodes PF4 Enhanced® assays					2		
ADAMTS13 Module							
ADAMTS13 (activity, inhibitor screen, titer, and anti-ADAMTS13 IgG)						3	

^{*}Not appropriate for meeting regulatory requirements; see page 166.

- CGS1, CGS2, CGS3 Two 2.0-mL lyophilized plasma specimens
- CGS4 Three 1.0-mL lyophilized plasma specimens
- CGS5 Two 60.0-µL serum specimens
- CGS7 Three 1.0-mL lyophilized plasma specimens in duplicate
- Two shipments per year

D-dimer Calibration Verification/Linearity LN42								
Analyte	Program Code							
	LN42	LN42 Target Range						
D-dimer	I	220-5,500 ng/mL FEU						

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL plasma specimens
- Two shipments per year

Fibrinogen Calibration Verification/Linearity LN44								
Analyte Program Code								
	LN44 LN44 Target Range							
Fibrinogen	I	80-900 mg/dL						

View your expedited linearity evaluations within two business days by logging into e-LAB Solutions Suite.

Program Information

- Six 1.0-mL frozen plasma specimens
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year; ships on dry ice

Coagulation Calibration Verification/Linearity LN35, LN36, LN37, LN51

Analyte		Progra	m Code		
	LN35	LN36	LN37	LN51	Target Ranges
Antithrombin activity					10%-130%
Protein C activity	I				10%-100%
Heparin, low molecular weight					0.1-2.0 U/mL
Heparin, unfractionated					0.1-1.3 U/mL
von Willebrand factor antigen					5%-140%
Factor VIII clot-based				ı	1-200 IU/dL
Factor VIII chromogenic				ı	1-200 IU/dL

View your expedited linearity evaluations for LN35, LN36, and LN37 within two business days by logging into e-LAB Solutions Suite.

The LN35, LN36, and LN37 CVL programs meet the CAP Accreditation Checklist requirements HEM.37363, 37365, 37373, and 37375.

- LN35, LN37, LN51 Six 1.0-mL frozen plasma specimens per mailing
- LN36 Twelve 1.0-mL frozen plasma specimens per mailing, which include six for low molecular weight heparin and six for unfractionated heparin
- Two shipments per year; ships on dry ice

Apixaban, Dabigatran, Fondaparinux, Rivaroxaban Anticoagulant Monitoring APXBN, DBGN, FNPX, RVBN

Analyte		Progra	Challenges per Shipment		
	APXBN	APXBN DBGN FNPX RVBN			
Activated partial thromboplastin time*		•			3
Prothrombin time*		I			3
Thrombin time		I			3
Apixaban					3
Dabigatran		I			3
Fondaparinux					3
Rivaroxaban					3

^{*}Not appropriate for meeting regulatory requirements; see page 166.

Program Information

- APXBN, DBGN, FNPX, RVBN - Three 1.0-mL lyophilized plasma specimens
- Two shipments per year

Activated Clotting Time Series CT, CT1, CT2, CT3, CT5						
Instrument/Cartridge		Pro	gram C	ode		Challenges per Shipment
	СТ	CT1	CT2	СТЗ	CT5	
Helena Actalyke C-ACT						3
Helena Actalyke MAX-ACT						3
IL GEM Hemochron 100/ACT+						3
IL GEM Hemochron 100/ACT-LR						3
IL Hemochron CA 510/FTCA510						3
IL Hemochron FTK-ACT						3
IL Hemochron P214/P215						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT+						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT-LR						3
i-STAT Celite® and Kaolin ACT					ı	3
Medtronic Hemotec ACT/ACTII/ACT Plus® HR-ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus LR-ACT		•				3
Medtronic Hemotec ACT/ACTII/ACT Plus R-ACT		•				3
Medtronic Hepcon HMS Plus						3

For multiple instrument reporting options, see the Quality Cross Check programs, CTQ, CT1Q, CT2Q, CT3Q, and CT5Q, on page 171.

- CT Three 3.0-mL lyophilized whole blood specimens with corresponding diluents
- CT1 Three 1.7-mL lyophilized whole blood specimens with corresponding diluents
- CT2 Three 0.5-mL lyophilized whole blood/ diluent ampules
- CT3 Three 0.5-mL lyophilized whole blood/ diluent ampules
- CT5 Three 1.7-mL lyophilized whole blood specimens with corresponding diluents
- · Two shipments per year

Quality Cross Check— Activated Clotting Time Series CTQ, CT1Q, CT2Q, CT3Q, CT5Q

Instrument/Cartridge		Pro		Challenges per Shipment		
	CTQ	CT1Q	CT2Q	CT3Q	CT5Q	
Helena Actalyke C-ACT®	ı					3
Helena Actalyke MAX-ACT						
IL GEM Hemochron 100/ACT+						
IL GEM Hemochron 100/ACT-LR						
IL Hemochron® CA510/FTCA510						3
IL Hemochron FTK-ACT						3
IL Hemochron P214/P215						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT+						3
IL Hemochron Signature Elite/ Hemochron Jr./ACT-LR						3
i-STAT Celite® and Kaolin ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus® HR-ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus LR-ACT						3
Medtronic Hemotec ACT/ACTII/ACT Plus R-ACT						3
Medtronic Hepcon HMS Plus						3

These programs do not meet regulatory requirements for proficiency testing; see programs CT-CT3 and CT5 on page 170. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics

Platelet I	unction	PF, PF1	
Instrument/Method	Progra	m Code	Challenges per Shipment
	PF	PF1	
Platelet aggregation			2
PFA-100, PFA-200			2
Helena Plateletworks®			2

These programs require the draw of a normal donor sample.

Program Information

- CTQ Three 3.0-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- CT1Q Three 1.7-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- CT2Q Three 0.5-mL lyophilized whole blood/ diluent ampules in triplicate
- CT3Q Three 0.5-mL lyophilized whole blood/ diluent ampules in triplicate
- CT5Q Three 1.7-mL lyophilized whole blood specimens in triplicate with corresponding diluents
- Report up to three instruments.
- Two shipments per year

- PF, PF1 Five 3.2% sodium citrate vacuum tubes; two 10.0-mL plastic tubes
- · Two shipments per year

Viscoelastic Studies VES							
Instrument	Program Code Challenges per Shipmen						
	VES						
TEG® 5000, TEG 6s, ROTEM® delta		2					

- Two 1.0-mL lyophilized plasma specimens
- · Two shipments per year

Viscoelastic Testing—Whole Blood VES1						
Instrument	Program Code	Challenges per Shipment				
	VES1					
Hemosonics Quantra®, ROTEM sigma, ROTEM delta	I	2				

This program requires the draw of a normal donor sample.

Program Information

- Four 3.2% sodium citrate vaccum tubes; two 4.0-mL pierceable cap tubes
- · Two shipments per year

Whole Blood Viscoelastic Assays in Clinical Diagnosis

The use of whole blood viscoelastic assays (TEG and ROTEM) to monitor or diagnose patients with various coagulopathies has increased exponentially in recent years. Whole Blood Viscoelastic Assays in Clinical

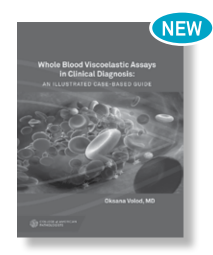
Diagnosis offers a practical and comprehensive case-based guide for tracings interpretation to ensure testing quality and patient safety. Pathologists and other laboratory personnel will all benefit from its case use studies for hemostatic disorders, pregnancy, trauma, cardiac surgery, and more.

Topics covered include:

- Overview of hemostasis physiology
- Viscoelastic testing
- Case studies addressing different hemostatic disorders
- Clinical uses of viscoelastic assays

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Item number: PUB231

Softcover; 342 pages; 2023

Drug-Specific Platelet Aggregation PIA/PIAX								
Procedure Program Code Challenges per Shipment								
	PIA	PIAX						
Aspirin assay			3					
PRU test			3					

- PIA Three lyophilized specimens with diluents
- PIAX All program PIA specimens in duplicate
- For use with the Accumetrics VerifyNow® System
- · Kit includes sufficient material to perform one assay; multiple assay reporting requires the purchase of PIAX.
- · Two shipments per year

Whole Blood Coagulation WP3, WP4, WP6, WP9, WP10							
Analyte	Challenges per Shipment						
	Program Code						
	WP3	WP4	WP6	WP9	WP10		
International normalized ratio (INR)	5 5 5 5 3						
Prothrombin time	5	5	5	5	_		

For method compatibility, see whole blood coagulation instrument matrix below.

Program Information

- WP3 Five 1.0-mL lyophilized plasma specimens with corresponding diluents
- WP4, WP6 Five 0.5-mL unitized lyophilized blood specimens
- WP9 Five 0.3-mL lyophilized plasma specimens
- Three shipments per year
- WP10 Three 0.3-mL lyophilized plasma specimens with corresponding diluents; two shipments per year

Whole Blood Coagulation, Instrument Matrix

Instrument Program Code					
	WP3	WP4	WP6	WP9	WP10
CoaguSense™					
IL GEM Hemochron Jr. Signature/Signature+, Signature Elite and Jr. II – citrated cuvette		•			
IL GEM Hemochron Jr. Signature/Signature+, Signature Elite and Jr. II – noncitrated cuvette					
i-STAT/i-STAT PTPlus					
Roche CoaguChek XS Plus, XS Pro, and CoaguChek Pro II					
Roche CoaguChek XS System					
Siemens Xprecia Stride					

Platelet Mapping PLTM							
Analyte	Program Code	Challenges per Shipment					
	PLTM						
AA % aggregation/inhibition	I	2					
ADP % aggregation/inhibition		2					

This program requires the draw of a normal donor sample.

Improve the reliability of your patient results with CAP Survey Validated Materials

Use the same material that is sent in the Surveys program to:

- Identify and troubleshoot instrument/method problems.
- · Correlate results with other laboratories or instruments.
- · Document correction of problems identified in Surveys.
- Utilize material with confirmed results as an alternative external quality control.
- · Identify potential proficiency testing failures.

Each laboratory receives a Survey Participant Summary, which includes readily available results.

Coagulation—Limited, Validated Material

Validated Material	Program Code	Corresponding Program	Page
Coagulation	CGM	CGL	166

Program Information

- One 3.2% sodium citrate and two heparin vacuum tubes; two 3.5-mL plastic tubes; one vial of 0.2M CaCl₂
- For use with the Haemonetics Platelet Mapping[®] assay
- · Two shipments per year

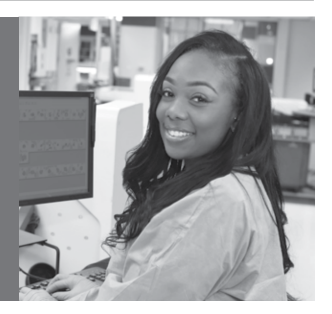
Program Information

 Seven 1.0-mL lyophilized plasma specimens and one 2.0-mL serum specimen; three shipments per year

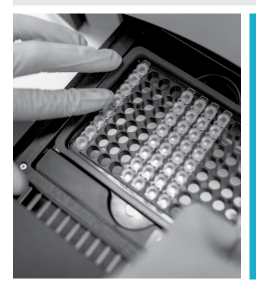
With direct transmission, less equals more.

Transmit your quantitative PT/EQA results directly to the CAP with direct transmission. Your laboratory will spend less time manually entering results, which will free up resources for other priorities. Plus, you will reduce clerical errors and streamline your process to be more like patient testing.

Get connected. Learn more at cap.org



15 Microbiology



Count on the CAP to support your PT/EQA needs in microbiology.

- New gastrointestinal panel for global laboratories (GIPN)
- Five-challenge PT/EQA programs for rapid malaria (RML5) and *Trichomonas vaginalis*, molecular (TVG5)

Microbiology

All current and new 2025 Centers for Medicare & Medicaid Services (CMS) regulated analytes are listed in **bold** type.

Microbiology	
Bacteriology	177
Mycobacteriology	193
Mycology	194
MycologyParasitology	197
Virology	199
Multidiscipline Microbiology	207
Infectious Disease Serology	
55	

New Programs NEW



Trichomonas vaginalis, Molecular, 5 Challenge (TVG5)	197
Rapid Malaria, 5 Challenge (RML5)	198
H5N1 Influenza A Detection and Subtyping (FLUA)	
Gastrointestinal Panel, Global (GIPN)	213

Discontinued Programs

licrobiolog

Microbiology

- Participants must report a minimum of five specimens, three times per year, to meet CLIA requirements for each of the subspecialties
 of microbiology (Bacteriology, Mycobacteriology*, Mycology, Parasitology, and Virology) for regulated testing.
 *Mycobacteriology requires five specimens, two times per year.
- · CLIA regulated tests are bolded.
- If any of the tests performed become waived by the FDA mid-year, your laboratory is responsible for maintaining five challenges per test event for the remaining non-waived tests in that subspecialty.

Guide to Molecular Microbiology Testing

Use this flowchart as a guide for ordering the appropriate Molecular Microbiology programs for your laboratory's testing menu. Participants must report five specimens for each mailing to meet CLIA requirements for the subspecialties of microbiology. See the following pages for more detailed information about each program.

Do you perform molecular testing on *Chlamydia* or GC only?

Do you perform nucleic acid amplification other than GC?

Do you perform viral load testing only?

Do you perform molecular multiplexing?

↓ YES

YES

↓ YES

↓ YES

Select from the following:

■ HC6, HC6X, HC7 Chlamydia/GC Nucleic Acid Amplification (page 191) Select from the following:

■ ID1, ID1T, ID2, ID5, IDN, IDO Nucleic Acid Amplification (pages 201, 204-205, 207)

■D1

Group A Streptococcus Culture/Molecular (page 179)

■ MRS2M, MRS5M MRSA Screen, Molecular (page 188)

BOR

Bordetella pertussis/ parapertussis (page 185)

■CDF5

C. difficile Detection (page 187)

MGEN

Mycoplasma genitalium (page 190)

■TVAG, TVG5

Trichomonas vaginalis
(page 197)

■VBDM

Zika (page 206)

■COV2, COVM SARS-CoV-2 (pages 202-203) Select from the following:

■HV2

HIV Viral Load (page 206)

■ HCV2, HBVL, HBVL5 Hepatitis Viral Load (page 205)

■VLS, VLS2 Viral Load (page 206) Select from the following:

■ID3

Nucleic Acid Amplification, Respiratory Limited (page 204)

■IDM5, IDME

Meningitis/Encephalitis Panel (page 209)

■IDPN

Infectious Disease Pneumonia Panel (page 211)

IDR

Infectious Disease Respiratory Panel (page 210)

■GIP, GIP5, GIPN

Gastrointestinal Panel (pages 212-213)

BCM

Bacterial Blood Culture (page 184)

MVP

Molecular Vaginal Panel (page 191)

■STIM

Sexually Transmitted Infection Detection (page 191)

Bacteriology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Guide for Ordering Regulated Bacteriology Programs

Procedure	Program Code							
	D D2 RMC D3 MC4 D1							
Bacterial identification	ı	I		ı		I		
Gram stain and morphology	ı	I		ı				
Antimicrobial susceptibility testing	ı	I						
Bacterial antigen/toxin detection	ı							

Participants must report five specimens for each mailing to meet CLIA requirements for the subspecialty of bacteriology. See the following pages for more detailed information about each program.

Bacteriology D			
Procedure	Program Code	Challenges per Shipment	
	D		
Antimicrobial susceptibility testing	I	2 graded	
Bacterial antigen/toxin detection		2	
Bacterial identification	I	5	
Gram stain and morphology	ı	1	

Additional Information

Antigen detection challenges will be included in the following shipments:

- Shipment A: C. difficile antigen/toxin and spinal fluid meningitis panel
- Shipment B: Spinal fluid meningitis panel, Group A Streptococcus, and C. difficile
- · Shipment C: C. difficile antigen/toxin and Group A Streptococcus

Program Information

- Five swab specimens with diluents in duplicate for culture
- Culture sources may include wounds, blood, respiratory, urines, stools, and anaerobes on a rotational basis.
- Three specimens for bacterial antigen/toxin detection from the following:

One swab for Group A Streptococcus

One 1.0-mL lyophilized specimen for spinal fluid meningitis testing

One 0.5-mL lyophilized specimen for *Clostridioides* (*Clostridium*) difficile, for use with rapid or molecular testing methods

· Three shipments per year







Refer to the Ordering Information provided for information regarding additional dangerous goods and related fees.

15

Expanded Bacteriology DEX				
Analyte	Program Code	Challenges per Shipment		
	DEX			
Bacterial identification	I	2		

Additional Information

Expanded Bacteriology (DEX) is an educational opportunity that provides:

- Culture and susceptibility testing challenges for microbiology laboratories that perform complete identification and susceptibility of bacterial isolates including less common or problematic bacteria
- More exposure to emerging bacterial pathogens and novel resistance mechanisms
- Ability to recognize and identify organisms that exhibit multiple drug-resistance patterns
- Recovery and identification of mixed pathogens such as yeast and bacteria (aerobic and anaerobic) in cultures containing multiple organisms

Program Information

- Two swab specimens in duplicate with diluents to perform bacterial identification and susceptibility (when directed)
- · Three shipments per year



Microbiology Bench Tools Competency MBT			
Procedure	Program Code	Challenges per Shipment	
	МВТ		
Bacterial identification		6	
Antimicrobial susceptibility testing	I	2	

Additional Information

Microbiology Bench Tools Competency (MBT) is a supplemental module for competency assessment and an educational resource for microbiology laboratories. The module:

- Provides organisms that challenge the basic elements of testing at the microbiology bench, including direct observation, monitoring, recording, and reporting of test results
- Can be used for both competency and educational purposes, including teaching and training pathology residents, new employees, and medical and MT/MLT students
- · Provides identification and susceptibility results for supervisor use

This is not a proficiency testing program and participants will not return results to the CAP.

Program Information

- Six swab specimens with diluents for bacterial identification and susceptibility
- Culture sources will vary with each shipment.
- Results will be provided with the kit to assess personnel competency.
- · Two shipments per year





Refer to the Ordering Information provided for information regarding additional dangerous goods and related fees.

Group A Streptococcus Culture/Molecular D1					
Procedure Program Code Challenges Shipmen					
	D1				
Bacterial identification	I	5			
Culture source:	Throat				
Microbiologic level:	Presence or absence of Group A Streptococcus determination				

- Five swab specimens with diluents in duplicate
- Throat swabs compatible with molecular- and culturebased methods
- Three shipments per year





GC and Urine Cultures D2, D3				
Procedure	Prograi	Program Code		
	D2	D2 D3		
Antimicrobial susceptibility testing			2	
Bacterial identification		1	5	
Gram stain and morphology		1	1	
Culture source:	Urine	Cervical		
Microbiologic level:	Organisms identified to the extent of your laboratory's protocol	Presence or absence of Neisseria gonorrhoeae determination		

Program Information

- D2 Five loop specimens with diluents in duplicate, with two susceptibility challenges and one Gram stain and morphology challenge
- D3 Five loop specimens with diluents in duplicate, and one Gram stain and morphology challenge
- Three shipments per year





Routine Microbiology Combination RMC				
Procedure	Program Code	Challenges per Shipment		
	RMC			
Antimicrobial susceptibility testing	I	2		
GC culture	I	2		
Gram stain and morphology	I	2		
Group A Streptococcus antigen detection*	I	1		
Throat culture/molecular	I	3		
Urine culture	I	3		

^{*}If your laboratory uses a waived method for Group A *Streptococcus*, these results will not count toward the required five challenges for the subspecialty of bacteriology.

- Five loop specimens with diluents in duplicate, three swab specimens with diluents in duplicate, and one swab specimen for bacterial antigen detection
- Urine culture will have two susceptibility challenges.
- Throat swabs compatible with molecular- and culturebased methods
- · Three shipments per year





Urine Colony Count	MC3, MC4	
Procedure	Challenges p	er Shipment
	Program Code	
	мсз	MC4
Urine colony count/urine culture identification	2	5
Group A Streptococcus antigen detection*		3
Throat culture/molecular		3

^{*}If your laboratory uses a waived method for Group A *Streptococcus*, these results will not count toward the required five challenges for the subspecialty of bacteriology.

Program Information

- MC3 Two urine specimens with diluents
- MC4 Five urine specimens with diluents, three swab specimens with diluents in duplicate, and three swab specimens for bacterial antigen detection
- Throat swabs compatible with molecular- and culturebased methods
- Three shipments per year





Gram Stain	D5	
Procedure	Program Code	Challenges per Shipment
	D5	
Gram stain and morphology		5

Program Information

- Five air-dried, methanolfixed, unstained glass slides
- · Three shipments per year





Assessment of Consistency of Gram Stain Morphologic Observations QPD10/QPD25

Introduction

Gram stain is a commonly performed bacterial stain in clinical microbiology laboratories. It is often the starting point guiding microbiological workup and initial clinical diagnosis and therapy. It is important for medical laboratory scientist/technologist staff who read Gram stains to provide an accurate interpretation based on reaction type and microscopic morphology, in order to provide presumptive identifications and quantification of bacteria and fungi in clinical specimens.

Objectives

This study will help assess the effectiveness of educational and practical experience policies and procedures dedicated to the laboratory's efforts in maintaining technologist skills in the morphological assessment of Gram stains. Participation in this study will help management assess the technologist's ability to evaluate Gram stains using online, whole slide images. These cases provide a standardized review and evaluation for each technologist. The study will help management meet applicable CLIA, CAP Laboratory Accreditation Program, and The Joint Commission laboratory requirements for morphology consistency of reporting among staff and personnel competency requirements (testing previously analyzed specimens).*

Data Collection

A series of online, whole slide images of Gram-stained smears using DigitalScope technology will be provided to each participating institution to assess technologists' ability to detect various microorganisms. Technologists will provide information about their work experience related to Gram stains, continuing education, and professional background. Information will be collected from each laboratory site about their continuing education requirements in microbiology and relevant laboratory procedures and policies related to Gram stain assessment. Each technologist will receive their own kit and result form.

Performance Indicators

- Individual technologist score (%) for each Gram stain case, and overall based on a standardized competency assessment method
- Overall laboratory score based on the facility's individual technologist performance(s)

Program Information

To meet your staff technical competency assessment requirements:

- Result forms for up to 10 technologists (QPD10)
- Result forms for up to 25 technologists (QPD25)
- Multiple orders may be purchased to accommodate the quantity of technologist result forms needed.

Preliminary study reports are provided at institution and technologist levels.

*Participation in this study helps laboratories meet:

- CLIA personnel requirements (Subpart M, 42 CFR §493.1)
- CAP Laboratory Accreditation Program Microbiology Checklist statements: MIC.11060, Culture Result Reporting, personnel performing
 Gram stains for this purpose are subject to competency assessment; MIC.11350, Morphologic Observation Evaluation, the laboratory
 evaluates consistency of morphologic observation among personnel performing microscopic analysis (eg, stains, wet preparations) from
 direct specimens and cultured organisms at least annually. The laboratory director or designee must determine acceptability criteria
 for agreement.
- CAP Laboratory Accreditation Program Checklist items: GEN.55500, element 5, Competency Assessment of Testing Personnel; GEN.55525, Performance Assessment of Supervisors/Consultants; DRA.11425, functions or responsibilities are properly performed by a qualified individual.
- The Joint Commission Standards HR.01.05.03, 01.06.01 (EPs 3, 18,19), HR.01.07.01, Pl.03.01.01(EPs 3-5), and LD.04.05.01, 04.05.03 (EPs 1-6) regarding in-service training, continuing education, competency, and evaluation of staff members

This is a one-time study conducted in the fourth quarter.

Virtual Gram Stain Competency VGS1, VGS2				
Procedure	Program Code Challenges per Shipment			
	VGS1	VGS2		
Virtual gram stain basic			3	
Virtual gram stain advanced	■ 3			

Additional Information

- Virtual Gram Stain Basic Competency (VGS1) is for general and new laboratory technologists/technicians. Participants will assess the quality of specimens and stains and will report artifacts and detailed gram-positive and gram-negative morphology. Challenges will include specimens such as CSF, body fluids, and positive blood cultures.
- Virtual Gram Stain Advanced Competency (VGS2) is for experienced laboratory technologists/technicians and microbiologists. Participants will receive challenging images of sputum, body fluids, and other specimens to assess the quality, quantity, and typical morphology of both gram-positive and gram-negative organisms appropriate for the site.
- See system requirements on page 12.

Program Information

- VGS1, VGS2 Three online whole slide images
- Results included in the kit to assess personnel competency
- Powered by DigitalScope® technology
- · Two shipments per year

Rapid Group A Strep Antigen Detection D6				
Procedure	dure Program Code Challenges per Shipmer			
	D6			
Group A Streptococcus antigen detection*	I	5		

^{*}If your laboratory uses a waived method for Group A Streptococcus, these results will not count toward the required five challenges for the subspecialty of bacteriology.

Program Information

- · Five swab specimens
- Not compatible with molecular- and culturebased methods
- · Three shipments per year



- Two swab specimens
- Not compatible with molecular- and culturebased methods
- · Two shipments per year

Group B Strep Detection D8					
Analyte Program Code Challenges per Shipme					
D8					
Group B Streptococcus ■ 5					

- Five swab specimens with diluents
- Program includes A549 cells to meet sample adequacy control requirement.
- Compatible with molecularand culture-based methods
- · Three shipments per year





Bacterial Antigen Detection LBAS, SBAS			
Procedure	Progra	m Code	Challenges per Shipment
	LBAS	SBAS	
Legionella pneumophila antigen detection	ı		2
Streptococcus pneumoniae antigen detection			2

Program Information

- LBAS, SBAS Two 0.5-mL liquid simulated clinical specimens
- Two shipments per year

Blood Culture BCS					
Procedure Program Code Challenges per Shipme					
BCS					
Blood culture bacterial and fungal detection and identification	ı	2			

Program Information

- Two specimens with diluents for inoculation of blood culture bottles
- Two shipments per year





Blood Culture, Staphylococcus aureus BCS1				
Analyte Program Code Challenges per Shipme				
	BCS1			
Staphylococcus aureus/MRSA	•	3		

Program Information

- Three specimens with diluents for inoculation of blood culture bottles
- Compatible with molecular methods for detection of S. aureus/MRSA from positive blood culture bottles
- Two shipments per year



Bacterial Blood Culture, Molecular BCM						
Procedure Program Code Challenges per Shipmer						
ВСМ						
Blood culture bacterial identification	re bacterial identification					

Additional Information

- This program is for the identification of gram-positive and gram-negative organisms, including common resistance mechanisms isolated from blood culture bottles.
- This program is not for the inoculation of blood culture bottles.

Program Information

- Five 1.0-mL simulated blood culture fluid specimens
- For laboratories using molecular multiplex panels
- · Three shipments per year

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Blood Culture Contamination QT2

Despite advances in blood culture practices and technology, false-positive blood culture results due to contaminants continue to be a critical problem. Blood culture contamination rate, the primary indicator of preanalytic performance in microbiology, is associated with increased length of hospital stay, additional expense, and the administration of unnecessary antibiotics.

The CAP and other accrediting organizations require you to monitor and evaluate key indicators of quality for improvement opportunities. Use this monitor to help meet CAP Laboratory Accreditation Program Checklist statements MIC.22630 and MIC.22635: "The laboratory must determine and regularly review the number of contaminated cultures. Tracking the contamination rate and providing feedback to units and persons drawing cultures is one method that has been shown to reduce contamination rates." This will also help laboratories meet The Joint Commission Standard QSA 04.07.01 EP3.

Objective

This study will determine the rate of blood culture contamination using standardized criteria for classifying contaminants.

Data Collection

On a monthly basis, participants will tabulate the total number of blood cultures processed and the total number of contaminated blood cultures. Blood cultures from neonatal patients are tabulated separately. For the purposes of this study, participants will consider a blood culture to be contaminated if they find one or more of the following organisms in only one of a series of blood culture specimens: Coagulase-negative *Staphylococcus*; *Micrococcus*; Alpha-hemolytic viridans group streptococci; *Propionibacterium acnes*; *Corynebacterium* sp. (diptheroids); or *Bacillus* sp. Participants have the option to monitor institution-specific subgroups (for example, a specific department or patient population).

Performance Indicators

- Neonatal contamination rate (%)
- Other contamination rate (%)
- Overall contamination rate (%)

Look in e-LAB Solutions Suite for your input forms approximately two weeks before the start of the next quarter.

Bordetella pertussis/parapertussis, Molecular BOR		
Analyte	Program Code	Challenges per Shipment
	BOR	
Bordetella pertussis	ı	3
Bordetella parapertussis	ı	3

- · Three swab specimens
- Designed for molecular techniques
- Two shipments per year

Carbapenemase Detection CRE		
Procedure	Program Code	Challenges per Shipment
	CRE	
Resistance mechanism detection		3

- Three swab specimens containing live organisms
- Designed for molecular and phenotypic testing methods
- Challenge isolates may include Enterobacterales, Pseudomonas, or Acinetobacter.
- Two shipments per year



Carbapenem-Resistant Organisms CRO		
Analyte	Program Code	Challenges per Shipment
	CRO	
KPC	ı	3
IMP	I	3
NDM	I	3
OXA-48	I	3
VIM	I	3

Program Information

- Three 130-µL specimens
- Designed for molecular techniques
- Compatible with Cepheid GeneXpert
- Two shipments per year

Campylobacter CAMP		
Analyte	Program Code	Challenges per Shipment
	CAMP	
Campylobacter	I	2

Program Information

- Two swabs with diluents in duplicate
- For use with rapid antigen, culture-based testing, and molecular methods
- Two shipments per year





C. difficile, 2 Challenge CDF2		
Analyte	Program Code	Challenges per Shipment
	CDF2	
Clostridioides (Clostridium) difficile antigen/toxin	ı	2

- Two 0.5-mL lyophilized specimens, for use with rapid or molecular testing methods
- · Two shipments per year

C. difficile, 5 Challenge CDF5		
Analyte Program Code Challenges per Shipmen		
	CDF5	
Clostridioides (Clostridium) difficile antigen/toxin	ı	5

Program Information

- Five 0.5-mL lyophilized specimens, for use with rapid or molecular testing methods
- · Three shipments per year

C. trachomatis Antigen Detection HC3		
Procedure	Program Code	Challenges per Shipment
	нсз	
C. trachomatis antigen detection (EIA)	I	5

Program Information

- Five 2.0-mL liquid specimens for Chlamydia antigen testing by EIA
- Three shipments per year

Fecal Lactoferrin FLAC		
Analyte	Program Code	Challenges per Shipment
	FLAC	
Fecal lactoferrin		3

Program Information

- Three 0.5-mL simulated stool specimens
- · For use with rapid methods
- Two shipments per year

Helicobacter pylori Antigen, Stool HPS				
Procedure	ure Program Code Challenges per Shipme			
	HPS			
Helicobacter pylori antigen		2		

Program Information

- Two 0.5-mL fecal suspensions
- Two shipments per year





Methicillin-Resistant <i>Staphylococcus aureus</i> Screen, 2 Challenge MRS			
Procedure Program Code Challenges per Shipmen			
MRS			
MRSA/MSSA detection		2	

Program Information

- Two swab specimens with diluents
- For laboratories performing culture-based testing only or using culture and molecular testing
- Two shipments per year



MRSA Screen, Molecular, 2 Challenge MRS2M		
Procedure Program Code Challenges per Shipment		
	MRS2M	
MRSA/MSSA/SA detection		2

Program Information

- Two swab specimens (in duplicate)
- For use with molecular methods that detect mecA
- Two shipments per year

Methicillin-Resistant <i>Staphylococcus aureus</i> Screen, 5 Challenge MRS5			
Procedure Program Code Challenges per Shipment			
	MRS5		
MRSA/MSSA detection	I	5	

Program Information

- Five swab specimens with diluents
- For laboratories performing culture-based testing only or using culture and molecular testing
- Three shipments per year



MRSA Screen, Molecular, 5 Challenge MRS5M					
Procedure	re Program Code Challenges per Shipment				
MRS5M					
MRSA/MSSA/SA detection	I	5			

Program Information

- Five swab specimens (in duplicate)
- For use with molecular methods that detect mecA
- Three shipments per year



Laboratory Preparedness Exercise LPX					
Analyte Program Code Challenges per Shipmen					
LPX					
Bacterial identification					

The Laboratory Preparedness Exercise (LPX) was developed as a collaborative effort between the College of American Pathologists, the Centers for Disease Control and Prevention (CDC), and the Association of Public Health Laboratories (APHL). Laboratories will be sent live organisms that either exhibit characteristics of bioterrorism agents or demonstrate epidemiologic importance, and will be expected to respond following Laboratory Response Network Sentinel Laboratory Guidelines if a bioterrorism agent is suspected. All agents provided are excluded from the CDC's select agent list. These may include strains of *Bacillus anthracis*, *Yersinia pestis*, *Francisella tularensis*, and *Brucella abortus* that have been modified and are safe for testing in a laboratory that contains a certified Class II Biological Safety Cabinet and is capable of handling Category A and B agents.

Program Information

- Three swab specimens with diluents
- Not available to customers outside the US due to US export law restrictions
- · Two shipments per year







Rapid Urease RUR			
Analyte	Program Code Challenges per Shipmen		
	RUR		
Urease	I	3	

Program Information

- Three simulated gastric biopsy specimens
- For use with methods such as CLOTEST®
- Two shipments per year

Stool Pathogen SP, SPN, SP1				
Analyte	Р	Program Code Challenges per Shipr		
	SP	SPN	SP1	
Adenovirus 40/41		•		2
C. difficile antigen/toxin		•		2
Rotavirus		•		2
Shiga toxin				2
Norovirus				1

Program Information

- SP Two 1.0-mL liquid specimens; for use with rapid or molecular testing methods; not available to customers outside the US due to US export law restrictions
- SPN Two 1.0-mL liquid specimens; for use with rapid or molecular testing methods; intended for laboratories outside the US
- SP1 One 1.0-mL liquid specimen compatible with molecular methods only
- · Two shipments per year



Shiga Toxin ST				
Analyte	Program Code Challenges per Shipment			
ST				
Shiga toxin	a toxin			

Program Information

- Two 0.5-mL liquid specimens
- For use with direct shiga toxin testing only; not compatible with culture methods, cytotoxicity assays, or PCR
- Not available to customers outside the US due to US export law restrictions
- · Two shipments per year

Bacterial Vaginosis BV				
Procedure	Program Code	Challenges per Shipment		
	BV			
Bacterial vaginosis detection	I	3		

Program Information

- Three 1.0-mL liquid specimens
- For OSOM® BVBlue users
- · Two shipments per year

Vaginitis Screen VS, VS1				
Analyte Program Code			Challenges per Shipment	
	VS*	VS1**		
Candida sp.			5	
Gardnerella vaginalis	I		5	
Trichomonas vaginalis ***	I		5	

^{*}The biohazard warning applies to program VS.

Program Information

 VS - Five swabs for DNA probe technology; BD Affirm™ VP III probe detection method; three shipments per year



 VS1 - Five swabs for methods such as Sekisui OSOM Trichomonas Rapid Test, Trichomonas vaginalis; three shipments per year

Mycoplasma genitalium, Molecular MGEN Analyte Program Code Challenges per Shipment MGEN Mycoplasma genitalium ■ 3

Program Information

- Three 1.0-mL liquid specimens
- Designed for molecular techniques
- Two shipments per year



^{**}Molecular users are encouraged to use *Trichomonas vaginalis*, Molecular (TVAG or TVG5), on page 197.

^{***}Trichomonas vaginalis is only reported to CMS for the VS program.

Molecular Vaginal Panel MVP					
Analyte	alyte Program Code Challenges per Shipme				
MVP					
Candida species group	I	5			
Candida krusei	I	5			
Candida glabrata ■ 5					
Trichomonas vaginalis					
Bacterial vaginosis	I	5			

- Five 1.0-mL liquid simulated vaginal specimens
- Designed for molecular methods such as BD MAX, Hologic, and Cepheid
- Three shipments per year

C. trachomatis and N. gonorrhoeae by NAA HC6, HC6X, HC7			
Procedure	Program Code Challenges per Shipmen		
	HC6*, HC6X*	HC7	
Nucleic acid amplification (NAA)			5
Nucleic acid amplification (NAA/DNA)			5

^{*}The biohazard warning applies to programs HC6 and HC6X.

Program Information

- HC6 Three swab specimens and two 1.0-mL liquid simulated urine specimens
- HC6X Three swab specimens and two 1.0-mL liquid simulated urine specimens in duplicate
- Three shipments per year



- HC7 Five 1.5-mL simulated body fluid specimens; designed for Cepheid users
- · Three shipments per year

Sexually Transmitted Infection Detection, Molecular STIM					
Analyte Program Code Challenges per Shipmer					
	STIM				
Chlamydia trachomatis	■ 5				
Neisseria gonorrhoeae	■ 5				
Mycoplasma genitalium		5			
Trichomonas vaginalis 5					

Program Information

- Five 2.0-mL simulated urogenital specimens
- Designed for molecular multiplex methods
- · Three shipments per year



Vaginitis Screen, Virtual Gram Stain VS2			
Procedure Program Code Challenges per Shipmer			
	VS2		
Interpretation of gram-stained vaginal smears	1	3	

See system requirements on page 12.

Program Information

- Three online whole slide images
- Powered by DigitalScope technology
- Two activities per year; your CAP shipping contact will be notified via email when the activity is available

Trichomonas vaginalis, Molecular TVAG, TVG5			
Analyte Program Code			
Challenges per Shipment			
	TVAG TVG5 NEW		
Trichomonas vaginalis	3	5	

Note: Only analytes in TVG5 will meet CMS requirements for parasite identification.

_		
Program	Informatio	r

- TVAG Three 1.5-mL liquid specimens; two shipments per year
- TVG5 Five 1.5-mL liquid specimens; three shipments per year

Vancomycin-Resistant Enterococcus VRE		
Procedure	Program Code	Challenges per Shipment
	VRE	
Vancomycin-resistant <i>Enterococcus</i> (VRE) detection		2

Program Information

- Two swabs with diluents
- For use with molecular methods and culture-based testing
- Two shipments per year





Mycobacteriology

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Mycobacteriology E		
Procedure	Program Code	Challenges per Shipment
	E	
Acid-fast smear	I	1
Antimycobacterial susceptibility testing	ı	1 graded, 1 ungraded
Mycobacterial identification*		5

^{*}This procedure requires identification of Mycobacterium tuberculosis.

Program Information

- Five simulated clinical isolates with diluents and one specimen for performing an acid-fast bacillus smear
- Identification may be performed by culture or molecular methods.
- · Two shipments per year





Mycobacteriology—Limited E1		
Procedure	Program Code	Challenges per Shipment
	E1	
Acid-fast smear		5
Mycobacterial culture		5

Program Information

- Five simulated specimens for acid-fast smears and/or for the determination of the presence or absence of acid-fast bacillus by culture
- · Two shipments per year



Molecular MTB Detection and Resistance MTR5, MTBR		
Procedure Challenges per Shipment		
Program Code		m Code
MTR5 MTBR		MTBR
Mycobacterium tuberculosis detection*	5	3
Rifampin resistance	5	3

^{*}Mycobacterium tuberculosis detection is only reported to CMS for the MTR5 program.

Program Information

- MTR5 Five 1.25-mL simulated sputum specimens for use with molecular methods
- MTBR Three 1.25-mL simulated sputum specimens for use with molecular methods
- Not suitable for culture
- Two shipments per year



Mycology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Mycology and Aerobic Actinomycetes F		
Procedure	Program Code	Challenges per Shipment
	F	
Antifungal susceptibility testing		1
Cryptococcal antigen detection		1
Mold and yeast identification		5

Program Information

- Five loops for culture with diluents in duplicate and one 1.0-mL simulated cerebrospinal fluid specimen
- Identification of yeasts, molds, and aerobic actinomycetes may be performed by molecular- and culture-based methods.
- · Three shipments per year





Yeast F1		
Procedure	Program Code	Challenges per Shipment
	F1	
Antifungal susceptibility testing	1	1
Cryptococcal antigen detection	1	1
Yeast identification	I	5

Program Information

- Five loops for culture with diluents in duplicate and one 1.0-mL simulated cerebrospinal fluid specimen
- Identification of yeast may be performed by molecularand culture-based methods.
- · Three shipments per year





Candida Culture F3		
Procedure	Program Code	Challenges per Shipment
	F3	
Yeast identification	1	5

Program Information

- Five loops for culture with diluents in duplicate
- · For laboratories identifying Candida sp. only
- Identification of Candida species may be performed by culture, molecular, and rapid methods.
- · Three shipments per year



Yeast Blood Culture, Molecular YBC		
Procedure	Program Code	Challenges per Shipment
	YBC	
Blood culture yeast identification		5

Additional Information

- This program is for identification of fungal organisms such as yeast isolated from blood culture bottles.
- This program is not for the inoculation of blood culture bottles.

Program In	nformation
------------	------------

- Five 1.0-mL simulated blood culture fluid specimens
- For laboratories using molecular multiplex panels
- · Three shipments per year

Cryptococcal Antigen Detection CRYP		
Procedure	Program Code	Challenges per Shipment
	CRYP	
Cryptococcal antigen	I	5

Cryptococcal Antigen Detection		CRYP
Procedure	Program Code	Challenges per Shipment
	CRYP	
Cryptococcal antigen	ı	5

Program Information

- · Five 1.0-mL simulated cerebrospinal fluids
- Three shipments per year

Galactomannan FGAL				
Analyte Program Code Challenges per Shipme				
	FGAL			
Galactomannan - Aspergillus	1	3		

Program Information

- Three liquid specimens
- For use with methods such as Bio-Rad Platelia™
- · Two shipments per year



Fungal Serology FSER				
Procedure	Program Code	Challenges per Shipment		
FSER				
Serological detection of specific fungal antibodies	ı	3		

- Three serum specimens
- For use with immunodiffusion methods
- Designed for the detection of IgG antibodies to Aspergillus, Blastomyces, Coccidioides, and Histoplasma
- Two shipments per year

Fungal Smear FSM				
Procedure	Program Code Challenges per Shipmen			
	FSM			
KOH preparation/calcofluor white	I	3		

Program Information

- Three unstained slides
- Two shipments per year

India Ink IND				
Procedure	Program Code	Challenges per Shipment		
IND				
India ink	I	2		

Program Information

- Two liquid specimens
- Two shipments per year

Pneumocystis jirovecii PCP1, PCP2, PCP4				
Procedure	Program Code Challenges per Shipmen			
	PCP1	PCP2	PCP4	
PCP – Calcofluor white stain	ı			3
PCP – DFA stain				3
PCP - GMS stain			ı	3

- PCP1, PCP2, PCP4 Three images, each available as photographs and online images for Pneumocystis jirovecii
- Two shipments per year

Parasitology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Parasitology P, P3, P4, P5				
Procedure	Procedure Challenges per Shipment			
	Program Code			
	Р	Р3	P4	P5
Fecal suspension (wet mount)	2	5	2	
Fecal suspension (Giardia and Cryptosporidium immunoassays and/or modified acid-fast stain)	2	1	1	5
Giemsa-stained blood smear	1			
Preserved slide (for permanent stain)	2		3	

Additional Information

- The proficiency testing materials used for the Parasitology programs contain formalin as a preservative.
- Number of specimen types are indicated in chart.

Program Information

- P Five specimens consisting of thin and thick films for blood and tissue parasite identification, preserved slides for permanent stain, 0.75-mL fecal suspensions for direct wet mount examination, photographs, and/or online images; two 0.75-mL fecal suspensions
- P3 Five 0.75-mL fecal suspensions for direct wet mount examination, photographs, and/or online images; one 0.75-mL fecal suspension
- P4 Five specimens consisting of 0.75-mL fecal suspensions for direct wet mount examination, preserved slides for permanent stain, photographs, and/or online images; one 0.75-mL fecal suspension
- P5 Five 0.75-mL fecal suspensions for Giardia and Cryptosporidium immunoassays and/or modified acid-fast stain
- P, P3, P4, P5 program specimens are for Giardia and Cryptosporidium immunoassays and/or modified acid-fast stain
- · Three shipments per year

Trichomonas vaginali	s, Molecular T\	/AG, TVG5
Analyte Program Code		
Challenges per Shipment		
	TVAG	TVG5 NEW
Trichomonas vaginalis	3	5

Note: Only analytes in TVG5 will meet CMS requirements for parasite identification.

- TVAG Three 1.5-mL liquid specimens; two shipments per year
- TVG5 Five 1.5-mL liquid specimens; three shipments per year

Blood Parasite BP			
Procedure	Program Code	Challenges per Shipment	
	ВР		
Blood parasite identification (thin/thick film sets*)	•	5	

^{*}This program will include corresponding thick films when available.

Program Information

- Five Giemsa-stained blood film sets, photographs, and/or online images
- Percent parasitemia reporting is provided when appropriate for educational purposes.
- A variety of blood parasites, including Plasmodium, Babesia, Trypanosoma, and filarial worms
- Three shipments per year

Rapid Malaria RMAL, RML5				
Analyte Program Code				
	Challenges per Shipment			
	RMAL	RML5 NEW		
Rapid malaria detection	3	5		
Plasmodium falciparum only	3	5		

Note: Only analytes in program RML5 will meet CMS requirements for parasite antigen detection.

Program Information

- RMAL Three 0.5-mL antigen specimens; two shipments per year
- RML5 Five 0.5-mL liquid specimens; three shipments per year

Expanded Parasitology PEX					
Procedure Program Code Challenges per Shipmer					
PEX					
Parasite identification	tification 3				

This program provides an educational opportunity to challenge laboratory professionals' competency in the identification of parasites utilizing photo images.

Program Information

- Three images, each available as photographs and online images
- · Two shipments per year

Ticks, Mites, and Other Arthropods TMO					
Procedure Program Code Challenges per Shipmer					
ТМО					
Tick, mite, and arthropod identification	ı	3			

Worm Identification WID					
Procedure	Program Code Challenges per Shipment				
WID					
Worm identification	■ 3				

Program Information

- Three images, each available as photographs and online images
- · Two shipments per year

- Three images, each available as photographs and online images
- · Two shipments per year

Virology

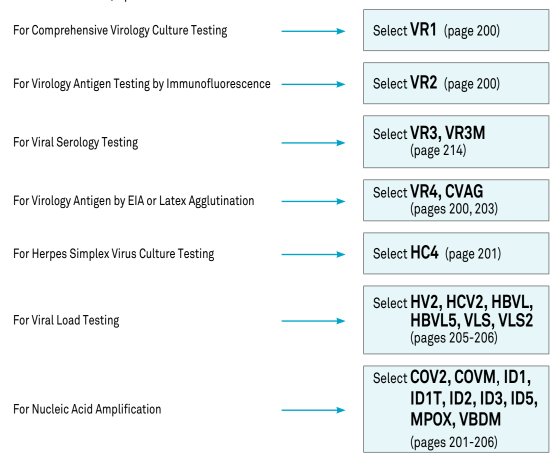
Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Guide for Ordering Regulated Virology Programs

Dragram Cada	Procedure		
Program Code	Viral Identification	Viral Antigen Detection	
VR1			
VR2			
VR4		ı	
HC4			
ID3			
ID5			
COVM			
CVAG			

Guide to Virology Testing

Use this flowchart as a guide for ordering the appropriate Virology programs for your laboratory's testing menu. For the subspecialty of virology, participants must test five specimens per mailing. If you have any questions, please call the Customer Contact Center at 800-323-4040 or +1-847-832-7000, Option 1.



Virology Culture VR1				
Procedure Program Code Challenges per Shipment				
	VR1			
Chlamydia trachomatis culture	I	1		
Viral isolation/identification ■ 5				

- Five 0.5-mL specimens for viral culture and one 0.5-mL specimen for Chlamydia trachomatis culture
- · Three shipments per year



Virology Antigen Detection (DFA) VR2				
Analyte/Procedure	Program Code	m Code Challenges per Shipment		
	VR2	Α	В	С
Adenovirus antigen		1	1	
Cytomegalovirus antigen		1	1	
Herpes simplex virus (HSV) antigen			1	1
Influenza A antigen		1		1
Influenza B antigen			1	
Parainfluenza antigen		1		1
Respiratory syncytial virus (RSV) antigen	•	1		1
Varicella-zoster (VZV) antigen			1	1
Educational challenge		1		

Program Information

- Five 5-well slide specimens
- Three shipments per year

Virology Antigen Detection (Non-DFA) VR4				
Analyte Program Code Challenges per Shipment				
	VR4			
Adenovirus (Not 40/41) antigen	ı	5		
Influenza A antigen	ı	5		
Influenza B antigen	I	5		
Respiratory syncytial virus (RSV) antigen	ı	5		
Rotavirus antigen	ı	5		

Program Information

- Five 1.5-mL specimens
- For use with enzyme immunoassay and/or latex agglutination methods
- Specimens not designed for molecular methods
- Three shipments per year



Herpes Simplex Virus HC4				
Procedure	Program Code Challenges per Shipment			
HC4				
Herpes simplex virus (HSV) culture ■ 5				

- Five 0.5-mL lyophilized specimens
- · Three shipments per year



Human Papillomavirus HPV			
Analyte	Program Code	Challenges per Shipment	
	HPV		
Human papillomavirus	I	2	

For laboratories using Digene, SurePath, and/or ThinPrep collection media, see page 308.

Program Information

- Two simulated cervical specimens contained in Digene transport media
- For Digene Hybrid Capture only
- Two shipments per year

Nucleic Acid Amplification, Viruses ID1, ID1T			
Analyte	Program Code		Challenges per Shipment
	ID1	ID1T	
Cytomegalovirus			1
Enterovirus	I		1
Epstein-Barr virus	I		1
Herpes simplex virus (HSV)			1
Human herpesvirus 6	I		1
Human herpesvirus 8	I		1
Parvovirus B19	I		1
Varicella-zoster virus (VZV)	I		1
BK virus			1
JC virus			1

Program Information

- ID1- Eight 1.0-mL liquid specimens
- ID1T Two 1.0-mL liquid specimens
- Two shipments per year



Mpox Molecular MPOX			
Procedure	Program Code Challenges per Shipment		
MPOX			
Monkeypox virus detection ■ 3			

This program is only available to customers within the US.

Program Information

- Three 1.0-mL simulated body fluid specimens that contain whole killed virus
- A549 cells included in each specimen
- For laboratories using molecular tests
- · Two shipments per year

SARS-CoV-2 Molecular COV2				
Analyte Program Code Challenges per Shipmer				
COV2				
SARS-CoV-2 ■ 3				

This program does not meet the proficiency testing requirements for laboratories subject to US Regulations and CAP-accredited laboratories that are performing non-waived testing. For multiple instrument reporting options, see the Quality Cross Check program, COV2Q, below.

Program Information

- Three 1.5-mL liquid simulated respiratory specimens
- Designed for molecular techniques
- Whole genome with sequence targets across all the assay platforms
- Qualitative and quantitative reporting options available
- · Two shipments per year

Quality Cross Check—SARS-CoV-2 Molecular COV2Q			
Analyte	Program Code Challenges per Shipment		
	COV2Q		
SARS-CoV-2		3	

This program does not meet regulatory requirements for proficiency testing; see program COV2, above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

- Three 3.2-mL non-infectious liquid specimens that contain the whole SARS-CoV-2 genome
- Designed for molecular techniques
- Report up to three instruments.
- Two shipments per year

SARS-CoV-2 Molecular, 5 Challenge COVM				
Analyte Program Code Challenges per Shipment				
соум				
SARS-CoV-2 ■ 5				

For multiple instrument reporting options, see the Quality Cross Check program, COV2Q, on page 202.

Program Information

- Five 1.5-mL liquid simulated respiratory specimens
- Designed for molecular techniques
- Whole genome with sequence targets across all the assay platforms
- Qualitative and quantitative reporting options available
- Three shipments per year

SARS-CoV-2 Antigen COVAG			
Analyte	Program Code Challenges per Shipmen		
	COVAG		
SARS-CoV-2 antigen	I	3	

This program does not meet the proficiency testing requirements for laboratories subject to US Regulations and CAP-accredited laboratories that are performing non-waived testing. For multiple instrument reporting options, see the Quality Cross Check program, COVAQ, below.

Program Information

- Three 0.5-mL simulated respiratory specimens
- · Designed for antigen test
- Two shipments per year

SARS-CoV-2 Antigen, 5 Challenge CVAG						
Analyte Program Code Challenges per Shipmer						
	CVAG					
SARS-CoV-2 antigen	■ 5					

For multiple instrument reporting options, see the Quality Cross Check program, COVAQ, below.

Program Information

- Five 0.5 mL simulated respiratory specimens
- · Designed for antigen test
- · Three shipments per year

Quality Cross Check—	ss Check—SARS-CoV-2 Antigen COVAQ				
Analyte	Program Code	Challenges per Shipment			
	COVAQ				
SARS-CoV-2 antigen	I	3			

This program does not meet regulatory requirements for proficiency testing; see program COVAG, above. For additional information about the Quality Cross Check program, see page 36.

Program Information Three 0.5-mL simulated

- respiratory specimens in triplicate
- Report up to three instruments.
- Two shipments per year

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

SARS-CoV-2 Serology COVS				
Analyte Program Code Challenges per Shipme				
covs				
SARS-CoV-2 antibody (total, IgG, IgM, and IgA) ■ 3				

- Three 0.5-mL serum specimens
- Appropriate for assays that detect antibodies to nucleocapsid, spike, combined antigen (nucleocapsid and spike), and the receptor binding domain of the spike protein
- Two shipments per year

Nucleic Acid Amplification, Respiratory ID2					
Analyte Program Code Challenges per Shipme					
	ID2				
Adenovirus	•	1			
Coronavirus/Rhinovirus*		1			
Human metapneumovirus		1			
Influenza virus*		1			
Parainfluenza virus		1			
Respiratory syncytial virus (RSV)					

- *Coronavirus/Rhinovirus and Influenza virus will be included in the following shipments:

 Shipment A: Coronavirus and Influenza A (does not include SARS-CoV-2)
 - Shipment B: Rhinovirus and Influenza B

Program Information

- Six 1.0-mL liquid specimens
- Two shipments per year

Nucleic Acid Amplification, Respiratory Limited ID3				
Analyte	Program Code	Challenges per Shipment		
	ID3			
Influenza A virus		5		
Influenza B virus		5		
Respiratory syncytial virus (RSV)		5		
SARS-CoV-2		5		

This program does not contain human genome material or sequences from human RNase P gene.

For multiple instrument reporting options, see the Quality Cross Check program, ID3Q, on page 205.

- Five 1.0-mL liquid specimens
- Designed for molecular multiplex panel users
- Three shipments per year

Quality Cross Check—Nucleic Acid Amplification, Respiratory Limited ID3Q						
Analyte Program Code Challenges per Shipmen						
	ID3Q					
Influenza A virus	ı	3				
Influenza B virus	1	3				
Respiratory syncytial virus (RSV)	ı	3				
SARS-CoV-2 ■ 3						

Additional Information

- This program does not contain human genome material or sequences from human RNase P gene.
- This program does not meet regulatory requirements for proficiency testing; see program ID3, on page 204. For additional information about the Quality Cross Check program, see page 36.

Program Information

- Three 1.0-mL liquid specimens
- Designed for molecular multiplex panel users
- Report up to three instruments.
- Two shipments per year

H5N1 Influenza and Subtypir	NEW		
Analyte/Procedure	e/Procedure Program Code		
	FLUA		
Influenza A detection		3	
Influenza A subtyping		3	

HSV, VZV—Molecular ID5						
Analyte Program Code Challenges per Shipment						
ID5						
Herpes simplex virus (HSV)	1	5				
Varicella-zoster virus (VZV)	ı	5				

Hepatitis Viral Load HCV2, HBVL, HBVL5				
Procedure	Cha	Challenges per Shipment		
	Program Code			
	HCV2	HBVL	HBVL5	
HCV genotyping	1			
HCV, qualitative	1			
HCV viral load	5			
HBV viral load		3	5	

Program Information

- Three 1.5-mL liquid specimens
- Includes Avian Influenza A (H5N1) and other seasonal Influenza A strains
- Two shipments per year

Program Information

- Five 1.0-mL liquid specimens
- Designed for molecular techniques
- Three shipments per year

- HCV2 Five 1.5-mL liquid plasma specimens; three shipments per year
- HBVL Three 1.5-mL plasma specimens; two shipments per year
- HBVL5 Five 1.5-mL plasma specimens; three shipments per year

HIV Viral Load HV2, HIVG				
Procedure Program Code Challenges per Shipmer				
	HV2	HIVG		
HIV-RNA viral load			5	
HIV genotyping*		I	1	

^{*}HIV genotyping is for laboratories reporting reverse transcriptase, protease, and/or integrase mutations.

Program Information

- HV2 Five 2.5-mL liquid specimens
- HIVG One 1.0-mL liquid specimen
- · Three shipments per year

Viral Load VLS, VLS2				
Procedure Program Code Challenges per Shipm				
	VLS VLS2			
BK viral load			2	
CMV viral load	I	•	2	
EBV viral load	I		2	
Adenovirus viral load			2	
HHV6 viral load			2	

Program Information

- VLS Six 1.0-mL EDTA plasma specimens; two shipments per year
- VLS2 Ten 2.0-mL EDTA plasma specimens; three shipments per year

Viral Load Calibration Verification/Linearity LN38, LN39, LN45, LN52					
Analyte Program Code					
	LN38	LN39	LN45	LN52	Target Ranges
CMV viral load	ı				316.0-8.0M IU/mL
HIV viral load					50.0-5.0M IU/mL
HCV viral load					50.0-280.0M IU/mL
HBV viral load					1.3 log-8.5 log IU/mL

View your expedited linearity evaluations for LN38, LN39, and LN45 within two business days by logging into e-LAB Solutions Suite

- LN38 Six 1.5-mL liquid plasma specimens
- LN39 Six 2.5-mL liquid plasma specimens
- LN45 Seven 2.5-mL frozen DNA specimens
- LN52 Seven 2.5-mL frozen DNA specimens
- Two shipments per year; LN45 and LN52 ship on dry ice

Vector-Borne Disease—Molecular VBDM			
Analyte	Program Code Challenges per Shipment		
	VBDM		
Zika virus		3	

- Three 1.5-mL liquid specimens
- Two shipments per year

Multidiscipline Microbiology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Guide for Ordering Regulated Molecular Multidiscipline Programs

Program Code	Procedure			
	Bacterial Identification	Viral Identification	Fungal Identification	Parasite Identification
IDR				
GIP5				
IDM5				
IDPN				

Nucleic Acid Amplification, Organisms IDO, IDN			
Analyte/Procedure	Program Code		Challenges per Shipment
	IDO	IDN	
Bordetella pertussis/parapertussis	1	ı	1
Legionella pneumophila/Chlamydia pneumoniae*	1	1	1
Methicillin-resistant Staphylococcus aureus	ı	I	1
Molecular typing (bacterial isolates)	•	ı	1
Mycobacterium tuberculosis	1		1
Mycoplasma pneumoniae	1	ı	1
Vancomycin-resistant Enterococcus	•		1

Program Information

- IDO Seven liquid or swab simulated clinical isolate specimens and two diluents
- IDN Six liquid or swab simulated clinical isolate specimens and two diluents; designed for international laboratories that cannot receive MTB
- Two shipments per year



- *Legionella pneumophila/Chlamydia pneumoniae will be included in the following shipments:
 - · Shipment A: Chlamydia pneumoniae
 - Shipment B: Legionella pneumophila

Joint Infection Panel JIP			
Analyte	Program Code	Challenges per Shipment	
	JIP		
Anaerococcus prevotii/vaginalis	1	5	
Bacteroides fragilis	1	5	
Candida albicans	1	5	
Citrobacter spp.	1	5	
Cutibacterium avidum/granulosum	1	5	
Enterobacter cloacae complex	ı	5	
Enterococcus faecalis	I	5	
Enterococcus faecium	1	5	
Escherichia coli	ı	5	
Finegoldia magna	I	5	
Haemophilus influenzae	•	5	
Kingella kingae	ı	5	
Klebsiella aerogenes	1	5	
Klebsiella pneumoniae group	1	5	
Morganella morganii		5	
Neisseria gonorrhoeae	1	5	
Parvimonas micra	1	5	
Peptoniphilus spp.		5	
Peptostreptococcus anaerobius	I	5	
Proteus spp.	1	5	
Pseudomonas aeruginosa		5	
Salmonella spp.	1	5	
Serratia marcescens	1	5	
Staphylococcus aureus	I	5	
Staphylococcus lugdunensis		5	
Streptococcus agalactiae	•	5	
Streptococcus pneumoniae		5	
Streptococcus pyogenes	1	5	

- Five 0.5-mL liquid specimens
- Designed for molecular multiplex panel users
- Program challenges may contain the following antimicrobial resistance genes on a rotational basis: CTX-M, IMP, KPC, mecA/C and MREJ, NDM, OXA-48like, vanA/B, and VIM.
- Three shipments per year

Meningitis/Encephalitis Panel IDME, IDM5			
Analyte	Challenges per Shipment		
	Program Code		
	IDME	IDM5	
Escherichia coli K1	3	5	
Haemophilus influenzae	3	5	
Listeria monocytogenes	3	5	
Neisseria meningitidis	3	5	
Streptococcus agalactiae	3	5	
Streptococcus pneumoniae	3	5	
Cytomegalovirus (CMV)	3	5	
Enterovirus	3	5	
Herpes simplex virus 1 (HSV-1)	3	5	
Herpes simplex virus 2 (HSV-2)	3	5	
Human herpesvirus 6 (HHV-6)	3	5	
Human parechovirus	3	5	
Varicella-zoster virus (VZV)	3	5	
Cryptococcus neoformans/gattii	3	5	

Note: Only IDM5 analytes in **bold** type will meet CMS requirements for bacteriology, fungal, and virology identification. For programs that include more than one subspecialty of microbiology, per CLIA, your laboratory is required to test five specimens, three times a year, for each subspecialty your laboratory performs.

Program Information

- IDME Three 1.0-mL liquid specimens; two shipments per year
- IDM5 Five 1.0-mL liquid specimens; three shipments per year
- Designed for molecular multiplex panel users

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Infectious Disease, Respiratory Panel IDR			
Analyte	Program Code	Challenges per Shipment	
	IDR		
Adenovirus		5	
Bocavirus		5	
Bordetella (pertussis, parapertussis, bronchiseptica, holmesii)	ı	5	
Chlamydia pneumoniae		5	
Coronavirus		5	
Human metapneumovirus		5	
Influenza A		5	
Influenza B		5	
Legionella pneumophila		5	
Mycoplasma pneumoniae		5	
Parainfluenza		5	
Respiratory syncytial virus (RSV)		5	
Rhinovirus/Enterovirus		5	
SARS-CoV-2*		5	

^{*}SARS-CoV-2 specimens do not contain human genome material or sequences from the human RNase P gene.

For programs that include more than one subspecialty of microbiology, per CLIA, your laboratory is required to test five specimens, three times a year, for each subspecialty your laboratory performs.

- Five 1.0-mL liquid specimens
- Designed for molecular multiplex panel users
- · Three shipments per year

Infectious Disease, Pneumonia Panel IDPN			
Analyte	Program Code	Challenges per Shipment	
	IDPN		
Acinetobacter calcoaceticus-baumannii complex	I	5	
Adenovirus	ı	5	
Coronavirus*	I	5	
Chlamydia pneumoniae	ı	5	
Enterobacter cloacae complex	I	5	
Escherichia coli	I	5	
Haemophilus influenzae	1	5	
Human metapneumovirus	•	5	
Rhinovirus/Enterovirus	•	5	
Influenza A	ı	5	
Influenza B		5	
Klebsiella aerogenes	•	5	
Klebsiella oxytoca	•	5	
Klebsiella pneumoniae group		5	
Legionella pneumophila	•	5	
Moraxella catarrhalis		5	
Mycoplasma pneumoniae		5	
Parainfluenza virus	•	5	
Proteus spp.	•	5	
Pseudomonas aeruginosa	•	5	
Respiratory syncytial virus (RSV)		5	
Serratia marcescens	•	5	
Staphylococcus aureus	•	5	
Streptococcus agalactiae	•	5	
Streptococcus pneumoniae	•	5	
Streptococcus pyogenes	•	5	

^{*}Laboratories performing SARS-CoV-2 testing, see the COVM/COV2 program on pages 202, 203. Includes antimicrobial resistance genes, as appropriate. For programs that include more than one subspecialty of microbiology, per CLIA, your laboratory is required to test five specimens, three times a year, for each subspecialty your laboratory performs.

- Five 1.0-mL liquid specimens
- Designed for molecular multiplex panel users
- Three shipments per year

dastromestinati anet an, an o			
Analyte	Challenges per Shipment		
	Program Code		
	GIP	GIP5	
Adenovirus	3	5	
Astrovirus	3	5	
Campylobacter	3	5	
Clostridioides (Clostridium) difficile, toxin A/B	3	5	
Cryptosporidium	3	5	
Cyclospora cayetanensis	3	5	
Entamoeba histolytica	3	5	
Enteroaggregative E. coli (EAEC)	3	5	
Enteropathogenic <i>E. coli</i> (EPEC)	3	5	
Enterotoxigenic <i>E. coli</i> (ETEC) LT/ST	3	5	
Escherichia coli 0157	3	5	
Giardia duodenalis (lamblia)	3	5	
Norovirus GI/GII	3	5	
Plesiomonas shigelloides	3	5	
Rotavirus A	3	5	
Salmonella	3	5	
Sapovirus	3	5	
Shiga-like toxin producing E. coli (STEC) stx1/stx2	3	5	
Shigella/Enteroinvasive E. coli (EIEC)	3	5	
Shigella	3	5	
Vibrio cholerae/Vibrio group	3	5	
Yersinia enterocolitica	3	5	

Gastrointestinal Panel GIP, GIP5

Note: Only GIP5 analytes in **bold** type will meet CMS requirements for bacteriology, parasitology, and virology identification. For programs that include more than one subspecialty of microbiology, per CLIA, your laboratory is required to test five specimens, three times a year, for each subspecialty your laboratory performs.

- GIP Three 1.0-mL simulated stool specimens; two shipments per year
- GIP5 Five 1.0-mL simulated stool specimens; three shipments per year
- Designed for molecular multiplex panel users
- Not available to customers outside the US due to US export law restrictions

Gastrointestinal Panel, Global GIPN			
Analyte	Program Code	Challenges per Shipment	
	GIPN		
Adenovirus	I	5	
Astrovirus	I	5	
Campylobacter	I	5	
Clostridiodes (Clostridium) difficile toxin A/B	ı	5	
Cryptosporidium	I	5	
Cyclospora cayetanensis	I	5	
Entamoeba histolytica	I	5	
Enteroaggregative E. coli (EAEC)	I	5	
Enteropathogenic E. coli (EPEC)	I	5	
Enterotoxigenic E. coli (ETEC) LT/ST	I	5	
Giardia duodenalis (lamblia)	I	5	
Norovirus GI/GII	I	5	
Plesiomonas shigelloides	I	5	
Rotavirus A	I	5	
Salmonella	I	5	
Sapovirus	I	5	
Shigella/Enteroinvasive E. coli (EIEC)	I	5	
Shigella	I	5	
Yersinia enterocolitica	I	5	

This program does not meet US CLIA regulatory requirements for proficiency testing. See program GIP5 on page 212.

- Five 1.0-mL simulated stool specimens
- Three shipments per year
- Intended for laboratories outside the US

Infectious Disease Serology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Infectious Disease Serology VR3, VR3M			
Analyte	Program Code		Challenges per Shipment
	VR3	VR3M	
Cytomegalovirus (CMV) – IgG, IgM, and total antibodies	I		1
Epstein-Barr virus (EBV) – VCA – IgG, IgM EBNA – IgG, IgM, and total antibodies EA – IgG	•		1
Helicobacter pylori – IgG, IgA, and total antibodies			1
Herpes simplex virus (HSV) – IgG antibody			1
Mycoplasma pneumoniae – IgG, IgM, and total antibodies			1
Mumps – IgG			1
Rubeola virus (English measles) – IgG antibody			1
Toxoplasma gondii — IgG, IgM, and total antibodies			1
Varicella-zoster virus (VZV) – IgG and total antibodies	I		1

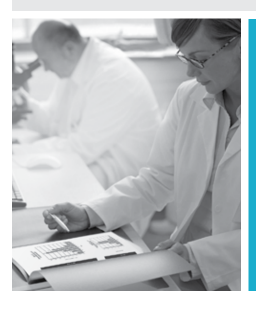
Program Information

- VR3 Eight 0.5-mL lyophilized defibrinated plasma specimens
- VR3M One 0.5-mL lyophilized defibrinated plasma specimen
- Two shipments per year

Tick-Transmitted Diseases TTD			
Analyte Program Code Challenges per Ship			
	TTD		
Antibodies to tick-transmitted disease organisms	•	3	

- Three 0.4-mL liquid specimens
- Designed for the detection of antibodies to Borrelia burgdorferi, Babesia microti, and Anaplasma phagocytophilum
- · Two shipments per year

16 Immunology and Flow Cytometry



Use the CAP's participant summaries to take your laboratory to the next level.

- Compare your results and methods against large peer groups for greater diagnostic confidence.
- Review the extensive discussion to further educate staff on testing trends and best practices.
- Earn continuing education credit with content that aligns with the proficiency testing challenge.

Immunology and Flow Cytometry

Immunology	216
Flow Cytometry	224

Discontinued Programs

Flow Cytometry—T-cell Subsets Analysis (FL7) Rare Flow Antigen Validation, CD103 (RFAV2)

Immunology

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Immunology ANA, ASO, CRP, HCG, IM, RF/RFX, RUB/RUBX, IL									
Analyte		Program Code				Challenges per Shipment			
	ANA	AS0	CRP	HCG	IM	RF/ RFX	RUB/ RUBX	IL	
Antinuclear antibody (ANA)*	•							ı	5
Antistreptolysin 0 (ASO)*									5
C-reactive protein, qualitative/quantitative								•	2
hCG, serum, qualitative/ quantitative								•	5
Infectious mononucleosis					•				5
Rheumatoid factor*									5
Rubella (IgG)*								ı	5

^{*}These CLIA-required analytes may be reported as qualitative, titer, or quantitative. The quantitative results are not reported to CMS.

Program Information

- ANA, RUB Five 0.5-mL serum specimens
- ANA Three online educational pattern interpretation challenges per year
- ASO, HCG, RF Five 1.0-mL serum specimens
- CRP Two 0.5-mL serum specimens; not appropriate for high-sensitivity CRP (hsCRP) methods
- IM Five 0.6-mL serum specimens
- RFX All program RF specimens in duplicate
- RUBX All program RUB specimens in duplicate
- IL All immunology specimens except RFX and RUBX
- Conventional and International System of Units (SI) reporting offered
- Three shipments per year



Immunology, General IG/IGX **Analyte Program Code** Challenges per Shipment IG/IGX Alpha-1 antitrypsin 5 Complement C3 5 **Complement C4** 5 Haptoglobin 5 5 **IgA** 5 IgE 5 IgG 5 5 Total kappa/lambda ratio

- IG Ten 1.0-mL serum specimens
- IGX All program IG specimens in duplicate
- Conventional and International System of Units (SI) reporting offered
- · Three shipments per year



Immunology, Special and <i>H. pylori</i> IgG Antibody S2, S4, S5				
Analyte		Program (Challenges per Shipment	
	S2 Special	S4 Special, Limited	S5 <i>H. pylori</i> IgG Antibody	
Anticentromere antibody	ı			2
Anti-DNA antibody double-stranded	ı	I		2
Antiglomerular basement membrane (GBM), IgG antibody	•			2
Antimitochondrial antibody	ı			2
Antineutrophil cytoplasmic antibody (ANCA, anti-MPO, anti-PR3)				2
Anti-RNP antibody	I			2
Anti-Ro52 antibody	ı			2
Anti-Ro60 antibody	ı			2
Anti-Sm antibody	I			2
Anti-Sm/RNP antibody	I			2
Antismooth muscle antibody	ı			2
Anti-SSA antibody	ı			2
Anti-SSB antibody	I			2
Anti-SSA/SSB antibody	I			2
Antithyroglobulin antibody	ı			2
Antithyroid peroxidase antibody/ Antithyroid microsomal antibody				2
Ceruloplasmin	ı	I		2
Haptoglobin	ı	ı		2
Helicobacter pylori, IgG antibody	ı	•	ı	2
IgD	ı	ı		2
IgG	ı	•		2
IgG subclass proteins	I	•		2
Prealbumin (transthyretin)	ı	•		2
Total kappa/lambda ratio	ı			2
Transferrin	ı			2

Program S2 is not appropriate for antimitochondrial antibody assays that are specific for the M2 antibody. Refer to program H on page 218.

Infectious Mononucleosis, Waived IMW					
Analyte Program Code Challenges per Shipment					
IMW					
Infectious mononucleosis, waived					

Program Information

- S2 Twenty-two (0.5- to 1.0-mL) serum specimens
- S4 Eight (0.5- to 1.0-mL) serum specimens
- S5 Two 1.0-mL serum specimens
- Two shipments per year



- Three 0.6-mL serum specimens
- Two shipments per year

Alpha-2-Macroglobulin A2MG				
Analyte Program Code Challenges per Shipment				
A2MG				
Alpha-2-macroglobulin	I	3		

Antichromatin Antibody ACA				
Analyte Program Code Challenges per Shipn				
	ACA			
Antichromatin antibody	I	3		

Antifilamentous Actin IgG Antibody FCN Analyte Program Code Challenges per Shipment FCN Antifilamentous actin (f-actin) IgG antibody 3

Antihistone Antibody AHT					
Analyte Program Code Challenges per Shipmen					
AHT					
Antihistone antibody	I	3			

Analyte	Program Code	Challenges per Shipment
	Н	
Antimitochondrial M2 antibody (AMA-M2)	•	2

Antimitochondrial M2 Antibody H

Autoimmune Gastritis Markers APC				
Analyte	Program Code	Challenges per Shipment		
	APC			
Antiparietal cell antibody	ı	2		
Anti-intrinsic factor antibody		2		

Program Information

- Three 0.5-mL serum specimens
- Two shipments per year

Program Information

- Three 0.5-mL serum specimens
- Two shipments per year

Program Information

- Three 0.5-mL serum specimens
- Two shipments per year

Program Information

- Three 0.5-mL serum specimens
- Two shipments per year



Program Information

- Two 1.0-mL serum specimens
- Two shipments per year

- Two 1.0-mL serum specimens
- Two shipments per year

Antiphospholipid Antibody ACL				
Analyte	Program Code	Challenges per Shipment		
	ACL			
Anticardiolipin antibody (polyclonal, lgG, lgM, and lgA)		3		
Beta-2-glycoprotein I (polyclonal, lgG, lgM, and lgA)		3		

- Three 0.5-mL lyophilized serum specimens
- Two shipments per year

Antiphosphatidylserine Antibody APS			
Analyte	Program Code	Challenges per Shipment	
	APS		
Anticardiolipin antibody (polyclonal, IgG, IgM, and IgA)	1	3	
Antiphosphatidylserine antibody (IgG, IgM, and IgA)	1	3	
Beta-2-glycoprotein I (polyclonal, IgG, IgM, and IgA)		3	
Antiphosphatidylserine/prothrombin antibody (aPS/PT)	1	3	

Program Information

- Three 0.5-mL lyophilized serum specimens
- Two shipments per year

Antiribosomal P Antibody ARP					
Analyte Program Code Challenges per Shipmen					
ARP					
Antiribosomal P antibody ■ 3					

Program Information

- Three 0.5-mL serum specimens
- Two shipments per year

Anti-Saccharomyces cerevisiae Antibody ASC Analyte **Program Code** Challenges per Shipment ASC Anti-Saccharomyces cerevisiae antibody 2 ı (lgG and lgA)

- Two 1.0-mL serum specimens
- · Two shipments per year

Celiac Serology CES/CESX				
Analyte	Progra	m Code	Challenges per Shipment	
	CES	CESX		
Antiendomysial antibody (IgA and IgG)	•	ı	3	
Antiendomysial antibody screen (IgA and IgG)	I	I	3	
Antigliadin antibody (IgA and IgG)	I	I	3	
Antideamidated gliadin peptide (DGP) antibody (IgA and IgG)	•		3	
Anti-DGP antibody screen (IgA and IgG)	I	I	3	
Antitissue transglutaminase (tTG) antibody (IgA and IgG)			3	
Anti-DGP and anti-tTG antibody screen (IgA and IgG)	•		3	

- CES Three 0.3-mL serum specimens
- CESX All program CES specimens in triplicate
- Two shipments per year

Cyclic Citrullinated Peptide Antibody (Anti-CCP) CCP			
Analyte	Program Code	Challenges per Shipment	
	ССР		
Anti-CCP	ı	2	
Rheumatoid factor isotypes (IgA, IgM, and IgG)	ı	2	

Cyclic Citrullinated Peptide Antibody (Anti-CCP) CCP			
Analyte	Program Code	Challenges per Shipment	
	ССР		
Anti-CCP		2	
Rheumatoid factor isotypes (IgA, IgM, and IgG)	•	2	

Cytokines	CTKN	
Analyte	Program Code	Challenges per Shipment
	CTKN	
Interleukin (IL)-1 beta	ı	3
IL-2	ı	3
IL-6	ı	3
IL-8	ı	3
IL-10	I	3
Tumor necrosis factor (TNF)-alpha	ı	3
Vascular endothelial growth factor (VEGF)	I	3

Program Information

- Two 1.0-mL serum specimens
- Two shipments per year



- Fifteen 1.0- to 3.0-mL lyophilized serum specimens
- Two shipments per year

Diagnostic Allergy SE			
Analyte/Procedure	Program Code	Challenges per Shipment	
	SE		
IgE, multiallergen screen, qualitative		5	
IgE, total		5	
Specific allergens		25	

- Five 2.0-mL serum specimens
- Includes common allergens from North America as well as less-frequently tested allergens
- Three shipments per year

High-Sensitivity C-Reactive Protein HSCRP		
Analyte	Program Code	Challenges per Shipment
	HSCRP	
High-sensitivity C-reactive protein		5

Program Information

- Five 0.5-mL liquid serum specimens
- Three shipments per year

Liver-Kidney Microsomal Antibody (Anti-LKM) LKM		
Analyte	Program Code	Challenges per Shipment
	LKM	
Anti-LKM		2

Program Information

- Two 0.3-mL serum specimens
- Two shipments per year

<i>M. tuberculosis</i> -Stimulated Infection Detection QF			
Analyte	te Program Code Challenges per Shipmer		
	QF		
M. tuberculosis		2	

This program is appropriate for the Autobio AutoLumo Series, QIAGEN QuantiFERON®-TB Gold and Gold Plus, DiaSorin Liaison QuantiFERON-TB Gold Plus, and SD Biosensor Standard methods.

Program	Information
• Two 1.0	0-mL lvophi

- Two 1.0-mL lyophilized serum specimens and one lyophilized mitogen control
- Two shipments per year

Rheumatic Disease Special Serologies RDS		
Analyte	Program Code	Challenges per Shipment
	RDS	
Anti-Jo-1 (antihistidyl t-RNA synthetase)		1
Anti-Scl-70 (anti-DNA topoisomerase)		1

- Two 1.0-mL serum specimens
- Two shipments per year



SARS-CoV-2 Serology COVS		
Analyte	Program Code	Challenges per Shipment
	covs	
SARS-CoV-2 antibody (total, IgG, IgM, and IgA)	I	3

- Three 0.5-mL serum specimens
- Appropriate for assays that detect antibodies to nucleocapsid, spike, combined antigen (nucleocapsid and spike), and the receptor binding domain of the spike protein
- · Two shipments per year

Syphilis Serology G		
Analyte	Program Code	Challenges per Shipment
	G	
Syphilis		5

Use with VDRL, RPR, MHA-TP/TP-PA/PK-TP/TPHA, EIA, CMIA, multiplex flow immunoassay, TP-LIA IgG, FTA-ABS, and USR methods. Laboratories performing syphilis serology on CSF specimens may also use this program.

Total Hemolytic Complement CH50		
Analyte	Program Code	Challenges per Shipment
	CH50	
Total hemolytic complement, 50% lysis	ı	2

Program Information

- Five 1.5-mL serum specimens
- Three shipments per year



- Two 0.5-mL lyophilized serum specimens
- · Two shipments per year

Viscosity V		
Analyte	Program Code	Challenges per Shipment
	V	
Viscosity	ı	2

- Two 10.0-mL serum specimens
- Two shipments per year

Serum Free Light Chains SFLC		
Analyte	Program Code	Challenges per Shipment
	SFLC	
Kappa serum free light chain		3
Lambda serum free light chain		3
Kappa/lambda serum free light chain ratio and ratio interpretation	ı	3

Program Information

- Three 1.0-mL serum specimens
- Two shipments per year

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Flow Cytometry

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Flow Cytometry FL, FL1, FL2				
Procedure	Pr	Program Code Challenges per Shipment		
	FL	FL1	FL2	
DNA content and cell cycle analysis	ı		I	3
Lymphocyte immunophenotyping	ı			3

These programs are not appropriate for hematology analyzers with monoclonal antibody analysis.

Program Information

- FL1 Three 1.5-mL whole blood specimens
- FL2 Three 1.1-mL specimens; two fixed cell line specimens and one calibrator for DNA content and cell cycle analysis
- FL All program FL1 and FL2 specimens
- · Three shipments per year

Flow Cytometry—Immunophenotypic Characterization of Leukemia/Lymphoma FL3				
Procedure Program Code Challenges per Shipmer				
FL3				
Leukemia/lymphoma	I	2		

Additional Information

- Program FL3 is suitable for laboratories that perform technical and interpretive components of leukemia/lymphoma specimens or laboratories that perform the technical component only. This program satisfies proficiency testing requirements for laboratories performing general analysis of leukemia/lymphoma specimens.
- · Laboratories that provide only interpretation (without technical component) should order program FL5.
- · This program has stability of two days or less. The CAP cannot guarantee performance or offer credits for orders placed for shipment outside of the US and Canada.

· ·
containing a cell li
blood mixture sim

Program Information

- Two 1.1-mL specimens ine/whole ulating leukemia/lymphoma with clinical histories and pertinent laboratory data; online images of tissue sections, bone marrow, and/ or peripheral blood smears as clinically relevant and/or available
- · Two shipments per year

Flow Cytometry, CD34+ FL4			
Analyte Program Code Challenges per Shipmer			
	FL4		
CD34+	I	2	

- Two 1.5-mL stabilized human CD34+ specimens
- Two shipments per year

Flow Cytometry, Interpretation Only FL5		
Procedure Program Code Challenges per Shipmer		
	FL5	
Flow cytometry, interpretation only of leukemia/lymphoma	ı	3

- Program FL5 is suitable for laboratories that provide only interpretation of flow data with technical component performed at an outside laboratory.
- This program may be ordered by laboratories that perform both technical and interpretation components and that are interested in obtaining additional interpretive material.

Program Information

- Three online cases
 consisting of gated dot
 plots, clinical histories, and
 pertinent laboratory data;
 online images of tissue
 sections, bone marrow, and/
 or peripheral blood smears
 as clinically relevant and/or
 available
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Flow Cytometry—Post-immunotherapy Analysis FL6		
Procedure	Program Code	Challenges per Shipment
	FL6	
Post-immunotherapy flow cytometry analysis	I	3

Additional Information

- Program FL6 is appropriate for laboratories that perform flow cytometry analysis on specimens from patients treated with immunotherapy regimens that cause immunophenotypic changes to normal and/or neoplastic cells. These include anti-CD20 (rituximab), anti-CD19 (CAR T19), and anti-CD38 therapies (daratumumab), among others.
- Participation in this program alone does not satisfy proficiency testing requirements for laboratories performing more general analysis of leukemia/ lymphoma specimens.

Program Information

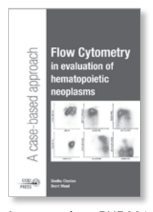
- Three online cases consisting of gated dot plots, clinical histories, and pertinent laboratory data
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Rely on this reference for a rapidly growing field.

Flow Cytometry in Evaluation of Hematopoietic Neoplasms: A Case-Based Approach is a practical guide to flow cytometric analysis in the workup of hematopoietic neoplasms presenting in the peripheral blood, marrow, lymphoid tissue, and extranodal sites.

Add it to your order, or view sample pages and purchase online.

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Item number: PUB221 Hardcover; 176 pages; 2012

Hematopathology Online Education HPATH/HPATH1		
Program	Program Code	Challenges per Shipment
	HPATH/HPATH1	
Hematopathology online case review	I	5

HPATH/HPATH1 prepares pathologists and pathology trainees with an interest in hematopathology to succeed by providing ongoing diagnostic learning and self-assessment.

The program includes 10 challenge cases per year written by expert hematopathologists. For each case the learner will:

- Review virtual whole slide image(s) in the context of a clinical scenario and relevant laboratory data.
- Select and interpret appropriate ancillary studies (IHC, flow cytometry, FISH, cytogenetics, molecular studies, etc) with real-time feedback.
- Simulate a real-world diagnostic workup to arrive at the appropriate diagnosis.
- Apply knowledge to answer three CME questions.

Cases span the breadth of benign and neoplastic hematopathology. Discussions incorporate current best practices in the workup, diagnosis, and subclassification of hematolymphoid diseases.

Neoplastic entities are discussed in the context of both the International Consensus Classification (ICC) and the fifth edition of the World Health Organization (WHO) classification.

See system requirements on page 12.

- HPATH Five diagnostic challenges per activity (two activities per year), with online whole slide images. Reporting with CME credit is available for one participant.
- HPATH1 Reporting option with CME credit for each additional participant (within the same institution); must order in conjunction with program HPATH
- Earn a maximum of 12.5 CME credits (AMA PRA Category 1 Credits™) per participant.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Powered by DigitalScope® technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



Flow Cytometry—B-ALL Measurable (Minimal) Residual Disease BALL		
Analyte	Program Code	Challenges per Shipment
	BALL	
B-ALL measurable (minimal) residual disease	ı	3

- Program BALL is intended for laboratories that perform measurable (minimal)
 residual disease (MRD) testing (rare event analysis) for B lymphoblastic leukemia/
 lymphoma. The cases presented will be a mixture of Children's Oncology Group
 (COG) approved B-ALL MRD method and laboratory developed assays.
- Participation in this program alone does not satisfy PT requirements for laboratories performing more general analysis of leukemia/lymphoma specimens.
- This program has stability of two days or less. The CAP cannot guarantee
 performance or offer credits for orders placed for shipment outside of the US and
 Canada.

Program Information

- Two 1.1-mL specimens containing a cell line/whole blood mixture simulating B lymphoblastic leukemia/ lymphoma measurable (minimal) residual disease
- One online case consisting of gated dot plots
- · Two shipments per year

Flow Cytometry—Mature B-cell Leukemia/Lymphoma Measurable (Minimal) Residual Disease FL8

Procedure	Program Code	Challenges per Shipment
	FL8	
Mature B-cell leukemia/lymphoma measurable (minimal) residual disease	ı	3

Additional Information

- Program FL8 is intended for laboratories that perform measurable (minimal) residual disease (MRD) testing (rare event analysis) for mature B-cell leukemia/ lymphoma.
- Participation in this program alone does not satisfy PT requirements for laboratories performing more general analysis of leukemia/lymphoma specimens.
- This program has stability of two days or less. The CAP cannot guarantee performance or offer credits for orders placed for shipment outside of the US and Canada.

- Two 1.1-mL specimens containing a cell line/whole blood mixture simulating mature B-cell leukemia/ lymphoma measurable (minimal) residual disease
- One online case consisting of gated dot plots
- · Two shipments per year

Flow Cytometry—Plasma Cell Myeloma Measurable (Minimal) Residual Disease FL9

Procedure	Program Code	Challenges per Shipment
	FL9	
Plasma cell myeloma measurable (minimal) residual disease	ı	3

Additional Information

- Program FL9 is intended for laboratories that perform measurable (minimal) residual disease (MRD) testing (rare event analysis) for plasma cell myeloma.
- Participation in this program alone does not satisfy PT requirements for laboratories performing more general analysis of leukemia/lymphoma specimens.
- This program has stability of two days or less. The CAP cannot guarantee
 performance or offer credits for orders placed for shipment outside of the US and
 Canada.

Program Information

- Two 4.5-mL specimens containing a cell line/whole blood mixture simulating plasma cell myeloma measurable (minimal) residual disease
- One online case consisting of gated dot plots
- · Two shipments per year

Flow Cytometry—Plasma Cell Neoplasms PCNEO				
Analyte	Program Code	Challenges per Shipment		
	PCNEO			
Plasma cell neoplasms	1	3		

Additional Information

- Program PCNEO is intended to supplement the FL3 program for laboratories
 performing both technical and interpretive components of leukemia/lymphoma
 analysis with specialized testing for plasma cells, including intracellular light chain
 (kappa/lambda) testing.
- Participation in this program alone does not satisfy PT requirements for laboratories performing more general analysis of leukemia/lymphoma specimens.
- This program has stability of two days or less. The CAP cannot guarantee
 performance or offer credits for orders placed for shipment outside of the US and
 Canada.

- One 1.1-mL specimen containing a cell line/whole blood mixture, simulating a plasma cell neoplasm with clinical history and pertinent laboratory data
- Two online cases consisting of gated dot plots, clinical histories, and pertinent laboratory data
- · Two shipments per year

Flow Cytometry—Immunophenotypic Characterization of Paroxysmal Nocturnal Hemoglobinuria PNH				
Analyte	Program Code Challenges per Shipment			
	PNH			
PNH RBC analysis	I	2		
PNH WBC analysis	I	2		

- The PNH program complies with the recommendations from the Guidelines for the Diagnosis and Monitoring of Paroxysmal Nocturnal Hemoglobinuria and Related Disorders by Flow Cytometry for RBC and WBC analysis. Due to the unique nature of these human donor-based materials, the shipping dates are subject to change. If this should occur, the CAP will provide notification prior to the originally scheduled shipping date.
- This program is appropriate for high-sensitivity testing (≤ 0.01% PNH type clone in red cells and/or granulocytes).

Fetal Red Cell Detection HBF Procedure Program Code Challenges per Shipment HBF Kleihauer-Betke and flow cytometry I 2 Rosette fetal screen I 2 Acid elution whole slide image I 1

Rare Flow Antigen Validation RFAV1, RFAV3 Analyte Program Code Challenges per Shipment RFAV1 RFAV3 CD1a 1 CD30 I 1

Additional Information

- Programs RFAV1 and RFAV3 do not meet the regulatory requirements for proficiency testing.
- These programs meet CAP Accreditation Checklist item FLO.23737, which requires semiannual testing of antigens.
- These programs have stability of two days or less. The CAP cannot guarantee
 performance or offer credits for orders placed for shipment outside of the US and
 Canada.

Program Information

- Two 0.5-mL whole blood specimens for RBC and WBC analysis
- Two shipments per year

Program Information

- Two 1.2-mL liquid whole blood specimens
- Not designed for F-cell quantitation
- Two online whole slide images per year with optional grids for cell counting
- Powered by DigitalScope technology
- Two shipments per year

- RFAV1 One 1.1-mL cell line specimen
- RFAV3 One 1.1-mL cell line specimen
- · Two shipments per year

ZAP-70/CD49d Analysis by Flow Cytometry ZAP70				
Analyte Program Code Challenges per Shipme				
	ZAP70			
Zeta-chain-associated protein kinase 70		3		
CD49d	I	3		

- This program tests for intracellular ZAP-70 staining of a cell line. It allows for assessment of the laboratory's staining techniques and the antibody clone used for ZAP-70 detection.
- CD49d is an important prognostic marker for CLL by flow cytometry. This program allows assessment of the laboratory's ability to detect CD49d.
- Laboratories may perform testing on ZAP-70, CD49d, or both.
- This program has stability of two days or less. The CAP cannot guarantee
 performance or offer credits for orders placed for shipment outside of the US and
 Canada.

Program Information

- Three 1.1-mL cell line specimens
- Two shipments per year

Color Atlas of Flow Cytometry

The Color Atlas of Flow Cytometry presents more than 70 cases from the CAP flow cytometry proficiency testing program, complete with over 270 images, photomicrographs, dot plots, survey data, and thorough discussions. Overviews of the hematopoietic disorders are also included with each section. Through peer-reviewed cases, practicing pathologists, medical technologists, residents, and students have an

opportunity to identify and appreciate disease categories and specific disease entities that are particularly difficult to diagnose correctly in clinical practice.

Topics include:

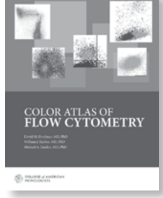
- B lymphoblastic leukemia and immature B cells
- Tlymphoblastic leukemia and immature T cells
- · Myeloid neoplasms
- Mature B-cell neoplasms

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All current and new 2025 Centers for Medicare & Medicaid Services (CMS) regulated analytes are listed in **bold** type.

Transfusion Medicine	232
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New Programs



Transfusion Medicine—Comprehensive (JXM)	. 232
Transfusion Medicine—Automated (JATXM)	

Discontinued Programs

Electronic Crossmatch (EXM), Electronic Crossmatch—Automated (EXM2) See Programs JXM and JATXM Bacterial Detection in Platelets, Rapid, 2 Challenge (BDPV) See Program BDPV5

Transfusion Medicine

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Transfusion Medicine J, JXM, J1, JE1					
Procedure		Program Code Challenges per Shipment			
	J	JXM NEW	J1	JE1	
ABO group	I	1			5
ABO subgroup					5
Rh typing	I		ı		5
Antibody detection	I				5
Antibody identification					5
Compatibility testing	I				5
Red blood cell antigen typing	I				1
Electronic crossmatch					3
Educational challenge					1

Program JXM assists laboratories in monitoring the performance of their electronic crossmatching system.

- J Five 3.0-mL 3% red blood cell suspensions; five 3.0-mL corresponding serum specimens; one 3.0mL donor red blood cell suspension
- JXM Five 3.0-mL 3% red blood cell suspensions; five 3.0-mL corresponding serum specimens; one 3.0mL donor red blood cell suspension; three simulated, ISBT 128 labeled donor unit challenges and three corresponding red blood cell suspensions
- J1 Five 3.0-mL 3% red blood cell suspensions; five 3.0-mL corresponding serum specimens
- JE1 One educational challenge, which may consist of a dry challenge and/or wet specimen for ABO grouping, ABO subgrouping, Rh typing, antibody detection, antibody identification, compatibility testing, antigen typing, and/ or direct antiglobulin testing
- Must order JE1 in conjunction with J or JXM programs.
- · Three shipments per year



Transfusion Medicine—Automated JAT, JATXM, JATE1				
Procedure	Progra	Program Code		Challenges per Shipment
	JAT	JATXM NEW	JATE1	
ABO group	•			5
ABO subgroup	•	ı		5
Rh typing	•	ı		5
Antibody detection	•			5
Antibody identification	•			5
Compatibility testing	•			5
Electronic crossmatch				3
Educational challenge				1

Program JATXM assists laboratories in monitoring the performance of their electronic crossmatching system.

Program Information

- JAT Five bar-coded 4.0-mL 13%-17% whole blood specimens and one 2.0-mL 23%-27% whole blood specimen for compatibility testing
- JATXM Five bar-coded
 4.0-mL 13%-17% whole
 blood specimens and
 one 2.0-mL 23%-27%
 whole blood specimen for
 compatibility testing; three
 simulated, ISBT 128 labeled
 donor unit challenges and
 three corresponding red
 blood cell suspensions
- JATE1 One educational challenge, which may consist of a dry challenge and/or wet specimen for ABO grouping, ABO subgrouping, Rh typing, antibody detection, antibody identification, and/or compatibility testing
- Must order JATE1 in conjunction with JAT or JATXM programs.
- · Three shipments per year



Quality Cross Check—Transfusion Medicine JATQProcedureProgram CodeChallenges per ShipmentJATQ3ABO grouping13Antibody detection13Rh typing13

This program does not meet regulatory requirements for proficiency testing; see program JAT above. For additional information about the Quality Cross Check program, see page 36.

The Quality Cross Check Program:

- Provides a solution for monitoring performance across multiple instruments and is in compliance with the CMS directive regarding proficiency testing on multiple instruments.
- Simplifies instrument comparability efforts by providing custom reports with both peer group comparison and instrument comparability statistics.

- Three 6.0-mL 13%-17% whole blood specimens
- May be used with automated and manual procedures
- Two shipments per year

In-Date Blood Product Wastage QT4

Blood for transfusion is a precious resource. At a minimum, wastage of blood that is not out-of-date represents a financial loss to the health care system. More ominously, systemic wastage of blood may reflect an environment of care that is out of control and may pose risks to patient safety.

Enrollment in this program assists laboratories in meeting regulatory requirements as follows:

- CAP Laboratory Accreditation Program Checklist statements TRM.40875, which requires the transfusion service medical director to monitor and audit transfusion practices to ensure the appropriate use of blood; TRM.30800, Disposition Records; and TRM.32275, Component Records, regarding recording the use of each blood or component product from receipt to final disposition.
- The Joint Commission Standards QSA.05.02.01, adequate blood and blood components; QSA.05.03.03, requirements for policies and procedures for returning unused blood products to blood transfusion services; and QSA.05.22.01, records of blood product disposition.
- AABB Standards for Blood Banks and Transfusion Services assessment 8.2, which requires transfusing facilities to have a peer-review program that monitors transfusion practices for blood components.

This study will compare the rates of blood product wastage (ie, units discarded in-date) in participating hospitals and track rates of improvement over time.

Data Collection

On a monthly basis, participants will use blood bank records to obtain information on the total number of units transfused for each type of blood component. Participants will track the number and type of blood units that are wasted in-date and the circumstances of wastage. This monitor includes the following types of blood components: whole blood (allogeneic), red blood cells (allogeneic), frozen plasma, platelet concentrates, single donor platelets, and cryoprecipitate.

Performance Indicators

- Overall blood wastage rate (%)
- Wastage rates by blood component type (%)

Performance Breakdown

• Breakdown of circumstances of wastage (%)

Look in e-LAB Solutions Suite for your input forms approximately two weeks before the start of the next quarter.

ABO Subgroup Typing ABOSG				
Procedure Program Code Challenges per Shipme				
	ABOSG			
ABO subgroup typing	I	3		
Rh typing	I	3		

ABO Subgroup Typing ABOSG					
Procedure Program Code Challenges per Shipmer					
	ABOSG				
ABO subgroup typing	I	3			
Rh typing	ı	3			

Red Blood Cell Antigen Genotyping RAG				
Procedure Program Code Challenges per Shipme				
	RAG			
RBC blood group genotyping for phenotype prediction	ı	3		

Red Blood Cell Antigen Typing RBCAT				
Procedure Program Code Challenges per Shipme				
	RBCAT			
Red blood cell antigen typing	I	2		

Program RBCAT is for donor centers and transfusion laboratories performing non-automated/manual red cell phenotyping for the management of patients with complex serology (ie, alloimmunization, sickle cell disease, warm autoimmune hemolytic anemia). Challenges will include antigens such as Rh, Kell, MNSs, Duffy, and Kidd blood group system.

Program Information

- Three 2.0-mL 3% red blood cell suspensions; three 2.0-mL corresponding serum specimens
- · Two shipments per year

Program Information

- Three 2.0-mL whole blood specimens
- · Two shipments per year

- Two 2.0-mL 2%-4% red blood cell suspensions
- · Two shipments per year

Antibody Titer ABT, ABT1, ABT2, ABT3						
Procedure			Progra	m Code		Challenges per Shipment
		ABT	ABT1	ABT2	ABT3	
Anti-A titer		I	ı			1
Anti-B titer						1
Anti-D titer						1

- ABT One 2.0-mL specimen for anti-A titer with one corresponding titer cell (3%-4% red blood cell suspension); one 2.0-mL specimen for anti-D titer with one corresponding titer cell (3%–4% red blood cell suspension)
- ABT1 One 2.0-mL specimen for anti-A titer with one corresponding titer cell (3%-4% red blood cell suspension)
- ABT2 One 2.0-mL specimen for anti-D titer with one corresponding titer cell (3%-4% red blood cell suspension)
- ABT3 One 2.0-mL specimen for anti-B titer with one corresponding titer cell (3%-4% red blood cell suspension)
- Two shipments per year

Antibody Titer—Automated AABT, AABT1, AABT2, AABT3					
Procedure	Program Code Challenges per Shipment				
	AABT	AABT1	AABT2	AABT3	
Anti-A titer		•			1
Anti-B titer					1
Anti-D titer	I		ı		1

- AABT One 2.0-mL specimen for anti-A titer; one 2.0-mL specimen for anti-D titer
- AABT1 One 2.0-mL specimen for anti-A titer
- AABT2 One 2.0-mL specimen for anti-D titer
- AABT3 One 2.0-mL specimen for anti-B titer
- Two shipments per year

Transfusion-Related Cell Count TRC					
Procedure Program Code Challenges per Shipment					
	TRC				
Platelet count (platelet-rich plasma)	ı	5			
WBC count		4			
Dry challenge		2			

WBC counts must be performed using a Nageotte chamber, by fluorescence microscopy, or by flow cytometry.

Direct Antiglobulin Testing DAT					
Procedure Program Code Challenges per Shipmen					
DAT					
Direct antiglobulin testing	Direct antiglobulin testing ■ 3				

Direct Antiglobulin Testing—Automated ADAT					
Procedure Program Code Challenges per Shipment					
	ADAT				
Direct antiglobulin testing	I	3			

Program Information

- Five 1.2-mL suspensions of platelet-rich plasma
- Two 1.0-mL vials leukocytereduced platelet material
- Two 1.0-mL vials leukocytereduced red blood cells
- Three shipments per year

Program Information

- Three 2.0-mL 3% red blood cell suspensions
- For use with manual method
- · Two shipments per year

- Three 4.0-mL 15% red blood cell suspensions
- For use with automated method
- Two shipments per year

Eluate Survey ELU				
Procedure Program Code Challenges per Shipme				
ELU				
Antibody elution 2				

Fetal Red Cell Detection HBF				
Procedure Program Code Challenges per Shi				
	HBF			
Kleihauer-Betke and flow cytometry	I	2		
Rosette fetal screen	I	2		
Acid elution whole slide image		1		

Program Information

• Two 2.0-mL 50% red blood cell suspensions • Two shipments per year

- Two 1.2-mL liquid whole blood specimens
- Not designed for F-cell quantitation
- Two online whole slide images per year with optional grids for cell counting
- · Powered by DigitalScope® technology
- Two shipments per year

Platelet Serology PS				
Procedure Program Code Challenges per Shipm				
	PS			
Antibody detection	I	3		
Platelet crossmatch	I	3		
Platelet antibody identification		3		

A low concentration of sodium azide may be present in the specimens and may affect lymphocytotoxicity methods.

- Three 3.0-mL plasma specimens
- · For use with solid-phase red cell adherence, flow cytometry, and EIA/ELISA methods
- Two shipments per year

Transfusion Medicine Comprehensive—Competency Assessment TMCA				
Procedure Program Code Challenges per Shipmen				
	TMCA			
ABO grouping		2		
Antibody detection		2		
Antibody identification		2		
Compatibility testing		2		
Rh typing		2		

Program TMCA does not meet the regulatory requirements for proficiency testing.

Program Information

or for an entire year

Program Information

• Two 3.0-mL 3% red blood cell suspensions

 Two 3.0-mL corresponding serum specimens
 One 3.0-mL donor 3% red blood cell suspension
 Three shipments per year; order shipments individually

- Two 2.0-mL 3% red blood cell suspensions
- Two shipments per year; order shipments individually or for an entire year

Direct Antiglobulin Test—Competency Assessment TMCAD					
Procedure Program Code Challenges per Shipment					
TMCAD					
Direct antiglobulin testing	Direct antiglobulin testing 2				

Program TMCAD does not meet the regulatory requirements for proficiency testing.

Eluate Competency Assessment TMCAE			
Procedure	ocedure Program Code		
	TMCAE		
Antibody elution	I	2	

Program TMCAE does not meet the regulatory requirements for proficiency testing.

Program Information

- Two 2.0-mL 50% red blood cell suspensions
- Two shipments per year; order shipments individually or for an entire year

Fetal Red Cell Quantitation—Competency Assessment TMCAF				
Procedure Program Code Challenges per Shipmen				
	TMCAF			
Kleihauer-Betke, flow cytometry	I	2		
Rosette fetal screen	I	2		
Acid elution whole slide image ■ 1				

Program TMCAF does not meet the regulatory requirements for proficiency testing.

- Two 1.2-mL whole blood specimens
- Two online whole slide images per year with optional grids for cell counting
- Powered by DigitalScope technology
- Two shipments per year; order shipments individually or for an entire year

Cord Blood and Stem Cell Processing CBT, SCP			
Analyte	Program Code		Challenges per Shipment
	СВТ	SCP	
Absolute CD3		ı	2
Absolute CD34			2
Bacterial culture	1	•	2
%CD3+		•	2
%CD34+	I	•	2
%CD45+		ı	2
CFU-GM	I	•	2
Total CFC	I	•	2
Fungal culture	1	ı	2
Hematocrit		•	2
Hemoglobin			2
Mononuclear cell count	1	ı	2
Nucleated red cells	I		2
Number of CD34 positive events			2
Number of CD45 positive events		•	2
Total nucleated cells	1	•	2
Viability	•	•	2
WBC count		ı	2

- Because these materials are human donor-based, the ship date is subject to change. If this should occur, notification will be provided prior to the scheduled date. In some instances, the program may ship in two installments.
- Due to material stability, no replacements will be available.
- These programs have stability of two days or less. The CAP cannot guarantee performance or offer credits for orders placed for shipment outside of the US and Canada.
- See International Shipping information section in the Ordering Information Supplement regarding additional dangerous goods shipping fees.

Program Information

- CBT Two 2.5-mL cord blood specimens; designed for assays required for the production of umbilical cord blood stem cell programs
- SCP Two 3.0-mL peripheral blood specimens; designed for laboratories that process and assess the suitability of stem cells
- · Two shipments per year





Refer to the Ordering Information provided for information regarding additional dangerous goods and related fees.

Bacterial Detection in Platelets BDP, BDP5			
Procedure	Program Code Challenges per Shipment		
	BDP	BDP5	
Bacterial culture and detection systems			2
Bacterial culture and detection systems			5

- The Centers for Medicare & Medicaid Services (CMS) requires proficiency testing for bacterial detection/identification. Please select the appropriate program for your laboratory based on the information below.
- Program BDP is designed for donor centers/laboratories that are associated with a CMS-certified microbiology laboratory with the same CLIA number, and which are participating in an approved proficiency testing program for bacterial detection.
- Program BDP5 is designed for donor centers/laboratories that are performing bacterial detection for the purposes of platelet unit screening and which are not associated with a CMS-certified microbiology laboratory with the same CLIA number.
- See International Shipping information section in the Ordering Information Supplement regarding additional dangerous goods shipping fees.

Program Information

- BDP Two lyophilized pellet specimens with diluents; two shipments per year
- BDP5 Five lyophilized pellet specimens with diluents; three shipments per year



Bacterial Detection in Platelets, Rapid BDPV5 Procedure Program Code Challenges per Shipment CMS certified rapid immunoassay BDPV5 5

Additional Information

- The Centers for Medicare & Medicaid Services (CMS) requires proficiency testing for bacterial detection in platelets.
- Program BDPV5 is designed for donor centers/laboratories that are performing bacterial detection for the purposes of platelet unit screening, and which are not associated with a CMS-certified microbiology laboratory with the same CLIA number.
- See International Shipping information section in the Ordering Information Supplement regarding additional dangerous goods shipping fees.

Program Information

- Five frozen specimens; three shipments per year
- For use with methods such as Verax Biomedical





Refer to the Ordering Information provided for information regarding additional dangerous goods and related fees.

Expanded Transfusion Medicine Exercises ETME1					
Procedure Program Code Challenges per Shipmen					
ETME1					
Expanded challenges	1	2			

Program ETME1 is an educational opportunity that offers:

- More challenging and/or complex antibody identification
- · Comprehensive case studies in transfusion medicine
- Simulated collaboration with other professionals, both those within and outside your institution
- A method for determining your laboratory's ability to recognize and integrate problem solving skills in transfusion medicine

The wet challenge may consist of specimens for ABO grouping, Rh typing, antibody detection, antibody identification, antigen typing, direct antiglobulin testing, and/or antibody elution.

Program Information

- One dry challenge and one wet challenge consisting of a serum specimen(s) and/or red blood cell suspensions
- · Two shipments per year

Transfusion Medicine: A Compendium of Educational Cases

Based on more than 10 years of educational material used in proficiency testing from the CAP Transfusion, Apheresis, and Cellular Therapy Committee, this newest book on transfusion medicine covers 20 cases with multiple-choice questions and answers. The topics included reflect clinical cases as well as hot topics

in transfusion medicine, and leverage the clinical experience of 19 highlyregarded transfusion medicine experts, all leaders in the field.

Contents include:

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Viral Markers

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Viral Markers—Series 1 VM1			
Analyte	Program Code	Challenges per Shipment	
	VM1		
Anti-HAV (total: IgM and IgG)	I	5	
Anti-HAV (IgG)	ı	5	
Anti-HBc (total: IgM and IgG)	ı	5	
Anti-HBs	ı	5	
Anti-HBs, quantitative	ı	5	
Anti-HCV	ı	5	
Anti-HIV-1	ı	5	
Anti-HIV-1/2	1	5	
Anti-HIV-2	ı	5	
HBsAg	ı	5	

Program Information

- Five 3.5-mL plasma specimens
- Three shipments per year

Additional Information

- Do not use program VM1 with rapid anti-HCV, anti-HIV-1, or anti-HIV-1/2 kits. See page 244 for programs appropriate for rapid methods.
- Anti-HIV-1/2, HIV-1 p24 antigen combination assay users should enroll in the VM6 program. Program VM1 is not appropriate for this assay.

Viral Markers—Series 2 VM2						
Analyte	yte Program Code Challenges per Shipme					
	VM2	VM2				
Anti-HBe	I	5				
HBeAg ■ 5						

Viral Markers—Series 3 VM3					
Analyte Program Code Challenges per Shipme					
VM3					
Anti-CMV	I 3				
Anti-HTLV-I/II	I 3				
HIV-1 p24 antigen ■ 3					

Program Information

- Five 3.5-mL plasma specimens
- · Three shipments per year

- Three 3.5-mL plasma specimens
- Two shipments per year

Viral Markers—Series 4 VM4				
Analyte	Program Code Challenges per Shipment			
	VM4			
Anti- <i>Trypanosoma cruzi</i> (Chagas disease)	I	2		

Viral Markers—Series 5 VM5 **Analyte Program Code** Challenges per Shipment VM5 Anti-HAV (IgM) 5 Anti-HBc (IgM) 5

Viral Markers—Series 6 VM6/VM6X **Program Code Analyte** Challenges per Shipment VM6 VM6X Anti-HIV-1/2 ı 5 5 HIV-1 p24 antigen

Anti-HIV 1/2 AHIV, AHIVW				
Analyte/Procedure Program Code Challenges per Shipmer				
	AHIV	AHIVW		
Anti-HIV-1, Anti-HIV-2, Anti-HIV-1/2			5	
Anti-HIV-1, Anti-HIV-1/2, waived methods only		•	2	

Anti-HCV, Rapid Methods, Waived RHCVW					
Analyte/Procedure Program Code Challenges per Shipn					
RHCVW					
Anti-HCV, waived methods only	I	3			

Program Information

- Two 1.0-mL plasma specimens
- Two shipments per year

Program Information

- Five 1.5-mL plasma specimens
- Three shipments per year

Program Information

- VM6 Five 0.5-mL plasma specimens
- VM6X All program VM6 specimens in duplicate
- Three shipments per year

Program Information

- AHIV Five 0.5-mL plasma specimens; three shipments per year
- AHIVW Two 0.5-mL plasma specimens; two shipments per year

- Three 0.5-mL plasma specimens
- Two shipments per year

Nucleic Acid Testing NAT			
Analyte	Program Code Challenges per Shipmen		
	NAT		
Babesia	I	1	
HBV	I	5	
HCV	I	5	
HIV	I	5	
West Nile virus	I	5	

- Five 6.0-mL plasma specimens
- One 1.1-mL whole blood specimen
- Designed for blood donor centers performing nucleic acid testing on donor units
- Compatible with HIV, HCV, and HBV multiplex assays
- Three shipments per year

Vector-Borne Disease—Molecular VBDM					
Analyte Program Code Challenges per Shipment					
VBDM					
Zika virus ■ 3					

Program Information

- Three 1.5-mL liquid specimens
- · Two shipments per year

Arthropod Benchtop Reference Guide

- Numerous identifications of ectoparasites commonly encountered in the clinical laboratory
- Detailed descriptions of the most significant morphologic elements, ecology, and clinical significance
- Eight tabbed sections for easy reference

 $\circ \ \, {\sf Introduction} \qquad \quad \circ \ \, {\sf Bed \ Bugs}$

• Ticks • Kissing Bugs

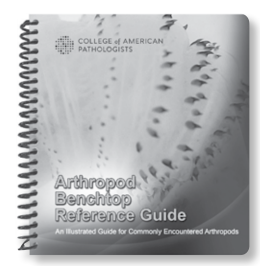
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Parentage Testing

Parentage/Relationship Test—Filter Paper PARF				
Analyte/Procedure Program Code Challenges per Shipm				
	PARF			
DNA testing (PCR)	I	4		
Calculation challenge (dry challenge) ■ 1				

Program Information

- DNA testing (PCR) Four samples per mailing: Two shipments of mother and child specimens on blood-stained filter paper with buccal swabs for two potential fathers; one shipment with all four specimens on blood-stained filter paper
- Reporting for short tandem repeats (STRs), X-STRs, Y-STRs, as well as the conclusions provided
- Three shipments per year

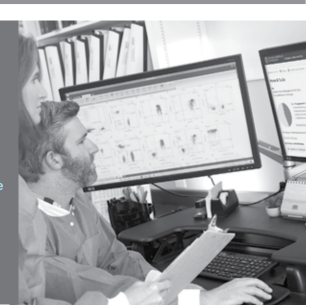
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18 Histocompatibility



Keep your laboratory current with insights from a panel of experts who monitor the latest trends in histocompatibility testing.

- Benefit from the CAP's culture of continuous improvement, which provides direction for updating our proficiency testing programs.
- Ensure your regulatory requirements are covered by continuing to participate in our programs.

New Programs

HLA Antibody Screen (Class I/Class II) Only (MXS)	248
HLA Crossmatching, Antibody Screen, and Antibody Identification (Class I/Class II),	
Extra Plasma (MXEP)	248

Program Changes

HLA Crossmatching, Antibody Screen, and Antibod	/ Identification (Class I/Class II)
Number of shipments and number of specimens	(MXC)248

Discontinued Programs

HLA Crossmatching, Antibody Screen, and Antibody Identification (Class I/Class II) (MXE) See Program MXEP

Histocompatibility

Antibody screen (Class I/Class II)

Antibody identification (Class I/Class II)

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

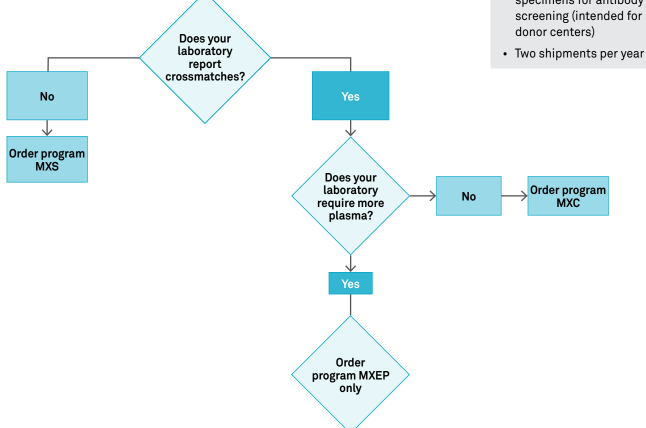
5

5

HLA Crossmatching, Antibody Screen, and **Antibody Identification (Class I/Class II)** MXC, MXEP, MXS Challenges per **Procedure Program Code Shipment MXC** MXS **MXEP** NEW NEW

5 Crossmatching (T-cell/B-cell) Program MXEP combines program MXC and the former program MXE, but with more plasma.

- MXC Five 0.4-mL plasma specimens; two (approximately 7–8 x 106 cells) purified blood lymphocyte specimens
- MXEP Five 0.4-mL plasma specimens in duplicate (0.8 mL total plasma); two (approximately 7–8 x 106 cells) purified blood lymphocyte specimens (intended for laboratories that require extra plasma volume for antibody identification)
- MXS Five 0.4-mL plasma specimens for antibody screening (intended for blood



Class I & II HLA Molecular Typing DML				
Procedure	Program Code	Challenges per Shipment		
	DML			
Molecular HLA-A, -B, and -C typing (Class I)	I	5		
Molecular HLA-DR, -DQ, and -DP typing (Class II)	ı	5		

- Five 2.0-mL whole blood specimens in CPD or CPD-A
- Serologic equivalents reporting available
- Two shipments per year

HLA-B27 Typing B27				
Procedure	Program Code	Challenges per Shipment		
	B27			
HLA-B27 typing	I	5		

Program Information

- Five 2.0-mL whole blood specimens in CPD or CPD-A
- Two shipments per year

Monitoring Engraftment ME				
Procedure	Program Code	Challenges per Shipment		
	ME			
Stem cell monitoring engraftment	1	5		

- Seven 0.5-mL whole blood specimens
- Designed for laboratories supporting stem cell transplant and laboratories monitoring chimerism after organ transplantation
- Two shipments per year

Antibody Titer ABT, ABT1, ABT2, ABT3					
Procedure		Progr	am Code	Challenges per Shipment	
	ABT	ABT1	ABT2	ABT3	
Anti-A titer		I			1
Anti-B titer					1
Anti-D titer	I		•		1

- ABT One 2.0-mL specimen for anti-A titer with one corresponding titer cell (3%-4% red blood cell suspension); one 2.0-mL specimen for anti-D titer with one corresponding titer cell (3%-4% red blood cell suspension)
- ABT1 One 2.0-mL specimen for anti-A titer with one corresponding titer cell (3%-4% red blood cell suspension)
- ABT2 One 2.0-mL specimen for anti-D titer with one corresponding titer cell (3%-4% red blood cell suspension)
- ABT3 One 2.0-mL specimen for anti-B titer with one corresponding titer cell (3%-4% red blood cell suspension)
- Two shipments per year

Antibody Titer—Automated AABT, AABT1, AABT2, AABT3					
Procedure	Program Code				Challenges per Shipment
	AABT	AABT1	AABT2	AABT3	
Anti-A titer		•			1
Anti-B titer				ı	1
Anti-D titer					1

- AABT One 2.0-mL specimen for anti-A titer; one 2.0-mL specimen for anti-D titer
- AABT1 One 2.0-mL specimen for anti-A titer
- AABT2 One 2.0-mL specimen for anti-D titer
- AABT3 One 2.0-mL specimen for anti-B titer
- · Two shipments per year

HLA Disease Assoc	iation—D	rug Risk	DADR1, DADR2
Analyte	Program Code		Challenges per Shipment
	DADR1	DADR2	
HLA-A*31:01			3
HLA-B*13:01			3
HLA-B*15:02			3
HLA-B*57:01			3
HLA-B*58:01			3
HLA-A*29:01			3
HLA-A*29:02			3
HLA-DQA1*04:01			3
HLA-DQA1*05:01		•	3
HLA-DQB1*03:02			3
HLA-DQB1*06:02			3
HLA-DRB1*03:01			3
HLA-DRB1*03:02		•	3
HLA-DRB1*04:02			3
HLA-DRB1*04:03			3
HLA-DRB1*04:06		•	3
HLA-DRB1*08:02			3
HLA-DRB1*08:04			3
HLA-DRB1*14:04			3
HLA-DRB1*14:05			3
HLA-DRB1*14:08			3
HLA-DRB1*15:01			3
HLA-DRB1*15:02			3
HLA-DQA1*02			3
HLA-DQA1*03			3
HLA-DQA1*05			3
HLA-DQB1*02:01			3
HLA-DQB1*02:02			3

These programs will challenge the laboratory to accurately identify the presence or absence of alleles associated with a variety of disease states (listed below) and/or the adverse reactions to specific drugs.

DADR1

- · Carbamazepine-induced Stevens-Johnson syndrome
- Allopurinol Stevens-Johnson syndrome
- Hypersensitivity to abacavir
- · Dapsone hypersensitivity

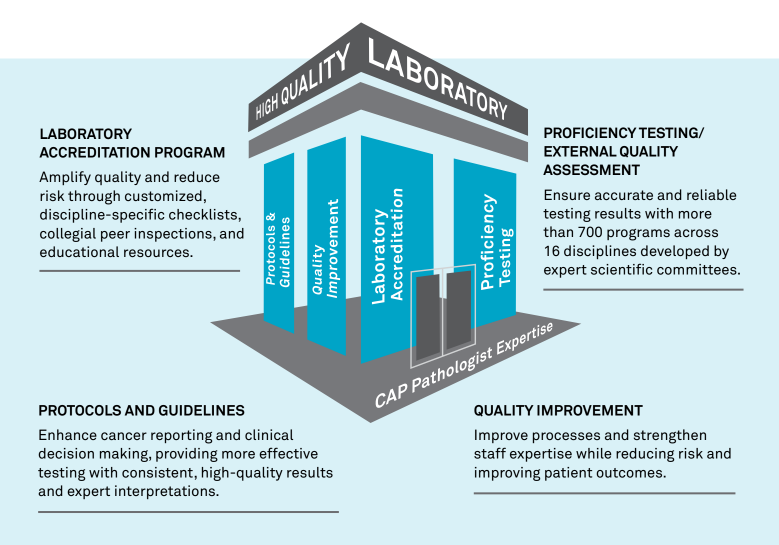
DADR2

- Celiac disease
- Narcolepsy
- · Pemphigus vulgaris
- Psoriasis
- Antiglomerular basement membrane disease
- · Birdshot retinochoroidopathy
- · Idiopathic myopathy

- DADR1, DADR2 Three 0.1-mL specimens, each containing 200 µg/mL of human DNA in media
- · Two shipments per year

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Genetics and Molecular Pathology



The CAP broadens its network of laboratory experts through its collaborations.

Among the organizations with which we partner:

- Association for Diagnostics & Laboratory Medicine (ADLM)
- American College of Medical Genetics and Genomics (ACMG)
- Association for Molecular Pathology (AMP)
- National Society for Histotechnology (NSH)

For additional information, refer to the CAP's Recommendations and Requirements for Molecular Proficiency Testing.

Genetics and Molecular Pathology

Cytogenetics	254
Biochemical and Molecular Genetics	
Next-Generation Sequencing	
Molecular Oncology—Solid Tumors	
Molecular Oncology—Hematologic	
Program Changes	
Next-Generation Sequencing—Solid Tumor (NGSST) additional	
paired normal specimen	266

Cytogenetics

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

CAP/ACMG Cytogenetics CY, CYBK				
Analyte/Procedure	Program Code Challenges per Shipmen			
	CY	СҮВК		
Karyotype abnormality			6	
Karyotype nomenclature			6	

Each challenge includes a case history and images of metaphase cells that are representative of each case. Each mailing will include three constitutional and three neoplastic challenges.

Program Information

- CY Online images of metaphase cells delivered two times a year; your CAP shipping contact will be notified via email when the activity is available
- CYBK Prints of metaphase cells; two shipments per year



CAP/ACMG Fluorescence In Situ Hybridization CYF, CYI				
Disease/Procedure	Program Code		Challenges per Shipment	
	CYF	CYI		
Constitutional and Hematologic Disorders				
FISH for constitutional disorder - slides			1	
FISH for constitutional disorder - dry challenge			2	
FISH for hematologic disorder - slides			1	
FISH for hematologic disorder - dry challenge			2	
Urothelial Carcinoma				
FISH for urothelial carcinoma			2	

Program Information

- CYF Four slides and four dry challenges
- CYI Two 250-µL cell samples suspended in ethanol from two different specimens; participants use FISH to detect chromosome abnormalities
- · Two shipments per year



Additional Information

• CYF 2025-A:

Constitutional disorder (two slides)-SRY Hematologic disorder (two slides)-20q del¹

• CYF 2025-B:

Constitutional disorder (two slides)-HIRA (TUPLE1) Hematologic disorder (two slides)-CBFB²

- ¹ For this challenge, participants should use the probe set used to interrogate deletion of 20q12 in their laboratories.
- ² For this challenge, participants should use the probe set used to interrogate CBFB rearrangements in their laboratories.
- CYF is prepared from cell suspension samples. For FISH in paraffin-embedded tissues, see page 255.
- These programs are only for laboratories that perform both hybridization and interpretation under the same CLIA number.

CAP/ACMG Fluorescence In Situ Hybridization for Paraffin-Embedded Tissue CYH, CYJ, CYK, CYL, CYALK

Analyte/Procedure	Program Code			Challenges per Shipment			
	СҮН	CYJ	СҮК	CYL	CYALK	Α	В
Breast Cancer							
ERBB2 (HER2) amplification						10	10
Interpretive challenges for ERBB2 (HER2) amplification						3	3
Brain/Glioma Tissue							
1p/19q						1	1
Solid Tumor							
ROS1 rearrangement						1	
DDIT3 (CHOP) rearrangement							1
Lymphoma Tissue							
CCND1 rearrangement						1	
ALK rearrangement							1
Lung Cancer							
ALK rearrangement					ı	1	
ALK rearrangement dry challenge					1		1

Program Information

- CYH Two unstained, fivecore tissue microarray slides equivalent to 10 paraffinembedded breast tissue specimens; two H&E stained tissue microarray slides are also provided
- CYJ Four unstained slides and one H&E stained slide
- CYK Two unstained slides and one H&E stained slide
- CYL Two unstained slides and one H&E stained slide
- CYALK Two unstained slides and one H&E stained slide are provided for the A mailing; the B mailing will include an ALK dry challenge.
- · Two shipments per year



Additional Information

- All CYJ, CYK, and CYL specimens will be 4.0-micron tissue sections mounted on positively charged glass slides.
- These programs are for laboratories that perform both hybridization and interpretation under the same CLIA number. For interpretation only ERBB2 (HER2) amplification by FISH for breast cancer, see program CYHI, below.

CAP/ACMG ERBB2 (HER2) Amplification by FISH, Interpretation Only CYHI Analyte/Procedure Program Code Challenges per Shipment CYHI ERBB2 (HER2) amplification in breast

Additional Information

cancer, interpretation only

- ERBB2 (HER2) Amplification by FISH, Interpretation Only is not considered proficiency testing. This exercise may be used to meet the requirements for alternative assessment.
- This program is for laboratories that perform <u>interpretation only</u> for *ERBB2 (HER2)* FISH for breast cancer.
- For laboratories that perform both hybridization and interpretation for *ERBB2* (*HER2*) FISH for breast cancer under the same CLIA number, see program CYH, above.

- Three online interpretation challenges; your CAP shipping contact will be notified via email when the activity is available
- · Two shipments per year



CAP/ACMG Constitutional Microarray CYCGH			
Procedure	Program Code	Challenges per Shipment	
	CYCGH		
Cytogenomic microarray analysis for constitutional abnormalities	ı	2	

- Participants will identify and characterize gains or losses and the cytogenetic location of abnormalities detected.
- This program is not appropriate for low resolution arrays that are designed to detect only aneuploidy.

CAP/ACMG Oncol	СҮСМА	
Procedure	Program Code	Challenges per Shipment
	СҮСМА	
Cytogenomic microarray analysis for oncologic abnormalities	ı	1

Participants will identify and characterize gains or losses and the cytogenetic location of abnormalities detected.

Program Information

- Two 2.0-µg DNA specimens
- Two shipments per year



- One 2.0-µg DNA specimen
- Two shipments per year



Biochemical and Molecular Genetics

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

CAP/ACMG Biochemical Genetics BGL, BGL1				
Analyte/Procedure	Progra	m Code	Challenges per Shipment	
	BGL	BGL1		
Acylcarnitines, qualitative and quantitative	•		1	
Amino acids, qualitative and quantitative	•		1	
Carnitine, qualitative and quantitative		•	3	
Glycosaminoglycans (mucopolysaccharides), qualitative and quantitative	1		1	
Organic acids, qualitative and quantitative	1		1	
Educational challenge			1	

Program Information

• BGL -

Acylcarnitines: One 0.1-mL plasma specimen

Amino acids: One 1.0-mL plasma or 2.0-mL urine specimen

Glycosaminoglycans (mucopolysaccharides): One 2.0-mL urine specimen

Organic acids: One 7.5-mL urine specimen

Educational challenge: Will consist of any one of the BGL analytes

- BGL1 Three 0.3-mL serum specimens
- · Two shipments per year



Sample Exchange Registry for Alternative Assessment

When no formal proficiency testing is yet available, join the CAP's Sample Exchange Registry. After at least three laboratories are identified as testing for the same rare analyte, the CAP can anonymously deliver a sample from each laboratory to another participating facility, all of whom then report their results to us. We send each participant a custom result report, including an anonymous participant summary covering all the laboratories that took part.

Learn more at cap.org

CAP/ACMG Amino Acid Quantitation for Inherited Metabolic Disorders BGL2

for innerited Metabolic Disorders BGL2				
Analyte/Procedure	Program Code	Challenges per Shipment		
	BGL2			
Alanine	I	3		
Alloisoleucine	I	3		
Arginine	I	3		
Aspartic acid	I	3		
Citrulline	I	3		
Cystine	I	3		
Glutamic acid	I	3		
Glutamine	I	3		
Glycine	I	3		
Histidine	I	3		
Homocystine	I	3		
Hydroxyproline	I	3		
Isoleucine	I	3		
Leucine	I	3		
Lysine	I	3		
Methionine	I	3		
Ornithine	I	3		
Phenylalanine	I	3		
Proline	I	3		
Serine	I	3		
Taurine	I	3		
Threonine	I	3		
Tryptophan	I	3		
Tyrosine	I	3		
Valine	I	3		

- Three 1.0-mL liquid specimens
- Two shipments per year



CAP/ACMG Acylcarnitine Quantitation for Inherited Metabolic Disorders BGL4 Analyte/Procedure **Program Code** Challenges per Shipment BGL4 3 Acetylcarnitine 3 Propionylcarnitine 3 Butyrylcarnitine Isovalerylcarnitine 3 Glutarylcarnitine 3 Hexanoylcarnitine 3 Octanoylcarnitine 3 Dodecanoylcarnitine Hexadecanoylcarnitine 3 3

Program Information

- Three 1.0-mL liquid specimens
- Two shipments per year



CAP/ACMG Alpha-1 Antitrypsin Genotyping AAT			
Analyte/Procedure	Program Code	Challenges per Shipment	
	AAT		
Alpha-1 antitrypsin (SERPINA1) genotyping		3	

This program will test for the M, S, and Z alleles.

3-OH-hexadecanoylcarnitine

Octadecanoylcarnitine

CAP/ACMG Apolipoprotein E Genotyping APOE				
Analyte/Procedure Program Code Challenges per Shipment				
	APOE			
Apolipoprotein E (APOE) genotyping	I	3		

This program is designed for laboratories utilizing APOE testing for hyperlipoproteinemia type III and Alzheimer diseases and will test for APOE e2, APOE e3, and APOE e4.

Program Information

3

- Three 10.0-µg extracted DNA specimens
- Two shipments per year



- Three 10.0-µg extracted DNA specimens
- Two shipments per year



CAP/ACMG BRCA1/2 Sequencing BRCA				
Analyte/Procedure	Program Code	Challenges per Shipment		
	BRCA			
BRCA1/2 DNA sequencing and variant interpretation	ı	3		
BRCA1/2 duplication/deletion analysis	ı	3		

Program Information

- Three 10.0-µg extracted **DNA** specimens
- · Two shipments per year



Additional Information

- Test your skill at reporting and interpreting DNA sequence variants for BRCA1/2 using standard nomenclature.
- Receive a summary and discussion of responses, including comments on the variant nomenclature and known or expected outcomes from identified variants.
- · Primers are not included; laboratories are expected to utilize the primers used in routine clinical testing.

CAP/ACMG Cardiomyopathy Sequencing Panel CMSP						
Analyte/Procedure	Analyte/Procedure Program Code Challenges per Shipment					
CMSP						
Cardiomyopathy sequencing panel	I	3				

Additional Information

- This proficiency challenge is for laboratories performing gene panels, exome sequencing, and whole genome sequencing to detect germline variants associated with inherited forms of cardiomyopathy.
- Participants will be asked to identify variants in the following genes: ACTC1, MYBPC3, MYH7, MYL2, MYL3, TNNI3, TNNT2, and TPM1.

- Three 80.0-µL purified extracted DNA specimens $(50 \text{ ng/}\mu\text{L})$
- · Two shipments per year



CAP/ACMG Hemoglobinopathies Genotyping HGM						
Analyte/Procedure Program Code Challenges per Shipment						
HGM						
Alpha-thalassemia	ı	3				
Beta-thalassemia	■ 3					
Hemoglobin S/C		3				

Program Information

- Three 50.0-µg extracted DNA specimens
- Two shipments per year



CAP/ACMG Inherited Cancer Sequencing Panel ICSP						
Analyte/Procedure Program Code Challenges per Shipment						
ICSP						
Inherited cancer sequencing panel	•	3				

Additional Information

- This proficiency challenge is for laboratories performing gene panels, exome sequencing, and whole genome sequencing to detect germline variants associated with inherited forms of cancer.
- Participants will be asked to identify variants in the following genes: APC, ATM, BRCA1, BRCA2, CDKN2A, CHEK2, MLH1, MSH2, MSH6, PALB2, and PMS2.

- Three 80.0-μL purified extracted DNA specimens (50 ng/μL)
- Two shipments per year



CAP/ACMG Molecular Genetics Series MGL1, MGL2, MGL3, MGL4, MGL5

	Program Code				O	
Disease/Gene	MGL1			MGL4	MGL5	Challenges per Shipment
Bloom syndrome (BLM gene)						3
BRCA1/2						3
Canavan (ASPA gene)						3
Connexin 26 (GJB2 gene)						3
Cystic fibrosis (CFTR gene)						3/2(MGL5)
DMD/Becker (<i>DMD</i> gene)						3
Factor V Leiden (F5 gene)	ı					3
Familial dysautonomia (ELP1 gene)						3
Fanconi anemia complementation group C (FANCC gene)				I		3
Fragile X (FMR1 gene)	ı					3
Friedreich ataxia (FXN gene)		•				3
Gaucher (GBA gene)						3
Glycogen storage disease type la (G6PC gene)				I		3
Hemochromatosis (HFE gene)	ı					3
Hemoglobin S/C						3
Huntington (HTT gene)						3
Methylenetetrahydrofolate reductase (MTHFR gene) c.665C>T (677C>T) and c.1286A>C (1298A>C)	•					3
Mucolipidosis IV (MCOLN1 gene)						3
Multiple endocrine neoplasia type 2 (<i>RET</i> gene)						3
Myotonic dystrophy (DMPK gene)						3
Niemann-Pick type A/B (SMPD1 gene)						3
Plasminogen activator inhibitor (PAI)-1 (SERPINE1 gene)	ı					3

Continued on the next page

Additional Information

- The BRCA1/2 program (module MGL3) is designed for laboratories testing for the three Ashkenazi Jewish founder mutations.
- The cystic fibrosis programs (modules MGL2 and MGL5) are designed for laboratories testing for the minimum mutation panel for population-based carrier screening from the ACMG Technical Standards and Guidelines for CFTR Mutation Testing, expanded panels, PolyT variant analysis, and/or full gene sequencing.
- Module MGL4 is designed for laboratories testing for diseases/disorders related to Ashkenazi Jewish ancestry.
- The Prader-Willi/Angelman syndrome program is designed for laboratories using methylation techniques for analysis.

- MGL1, MGL2, MGL3, MGL4 - Three 50.0-µg extracted DNA specimens per disease/gene
- MGL5 Two 50.0-µg extracted DNA specimens
- Two shipments per year



CAP/ACMG Molecular Genetics Series MGL1, MGL2, MGL3, MGL4, MGL5 continued						
Program Code Challenge					Challenges per	
Disease/Gene	MGL1	MGL2	MGL3	MGL4	MGL5	Shipment
Prader-Willi/Angelman syndrome	I					3
Prothrombin (F2 gene)						3
RhD		ı				3
Spinal muscular atrophy (SMN1 and SMN2 genes)						3
Spinocerebellar ataxia (ATXN1, ATXN2, ATXN3, CACNA1A, and ATXN7 genes)						3
Tay-Sachs (<i>HEXA</i> gene)						3

- The BRCA1/2 program (module MGL3) is designed for laboratories testing for the three Ashkenazi Jewish founder mutations.
- The cystic fibrosis programs (modules MGL2 and MGL5) are designed for laboratories testing for the minimum mutation panel for population-based carrier screening from the ACMG Technical Standards and Guidelines for CFTR Mutation Testing, expanded panels, PolyT variant analysis, and/or full gene sequencing.
- Module MGL4 is designed for laboratories testing for diseases/disorders related to Ashkenazi Jewish ancestry.
- The Prader-Willi/Angelman syndrome program is designed for laboratories using methylation techniques for analysis.
- The Spinal Muscular Atrophy program includes SMN1 and SMN2 gene analysis and copy number analysis.

CAP/ACMG Inherited Metabolic Diseases IMD1, IMD2, IMD3						
Analyte/Procedure	Program Code Challenges per Shipme					
	IMD1 IMD2 IMD3					
Mitochondrial DNA deletion syndromes				3		
MCAD				3		
Mitochondrial cytopathies*			ı	3		

^{*}Includes disorders/diseases such as Leber hereditary optic neuropathy and myoclonus epilepsy with ragged red fibers (MERRF).

Program Information

- MGL1, MGL2, MGL3, MGL4 - Three 50.0-µg extracted DNA specimens per disease/gene
- MGL5 Two 50.0-μg extracted DNA specimens
- · Two shipments per year



- IMD1, IMD2, IMD3 Three 50.0-µg extracted DNA specimens
- · Two shipments per year



CAP/ACMG Molecular Genetics Sequencing SEC, SEC1					
Procedure	Program Code Challenges per Shipment				
	SEC SEC1				
DNA sequencing interpretation challenge	■ 3				
DNA sequencing		•	3		

- Test your skill at interpreting and reporting DNA sequence variants for inherited diseases using standard nomenclature.
- Receive a summary and discussion of responses, including comments on nomenclature, known or expected outcomes from identified variants, and teaching points about genes/disorders represented.

Program Information

- SEC DNA sequence electropherogram files with a range of variants, suitable for base-calling and analysis using a range of commercial or public domain software programs; also includes nomenclature/variant references. Two online activities per year; your CAP shipping contact will be notified via email when the activity is available
- SEC1 Three 30.0-µg
 extracted DNA specimens;
 forward and reverse
 lyophilized primers are
 provided; two shipments
 per year



Pharmacogenetics PGX, PGX1, PGX3					
Analyte/Procedure	Program Code			Challenges per Shipment	
	PGX	PGX1	PGX3		
CYP2C19	ı			3	
CYP2C9	ı			3	
CYP2B6	ı			3	
CYP2D6	1			3	
CYP3A4	ı			3	
CYP3A5	ı			3	
CYP4F2	ı			3	
SLC01B1 (rs4149056)	ı			3	
VKORC1	ı			3	
IL28B (rs12979860)		ı		3	
COMT (rs4680)		I		3	
G6PD		I		3	
OPRM1 (rs1799971, c.118A>G)		ı		3	
DPYD				3	
NUDT15			ı	3	
TPMT				3	
UGT1A1				3	

UGT1A1 (PGX3 program) tests the laboratory's ability to detect variants in the TATA repeat sequence in the UGT1A1 promotor (eg, UGT1A1*28 with seven TA repeats). The ability to detect variants in other regions of the UGT1A1 gene is not part of this program.

- PGX, PGX1, PGX3 Three 25.0-µg extracted DNA specimens
- Includes allele detection (genotyping) and/or interpretive challenges
- Two shipments per year

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CAP/ACMG Rett Syndrome (MECP2) RETT					
Analyte/Procedure Program Code Challenges per Shipm					
	RETT				
Rett (<i>MECP2</i>) genotyping		3			
Rett (MECP2) duplication/deletion analysis	ı	3			

Program Information

- Three 10.0-µg extracted DNA specimens
- Two shipments per year



CAP/ACMG Thrombophilia Mutations TPM						
Analyte/Procedure Program Code Challenges per Shipment						
ТРМ						
Factor II (F2 gene, Prothrombin)	I	3				
Factor V Leiden (F5 gene)		3				

This program is designed for the Cepheid GeneXpert factor II and factor V assays. DNA extraction for other assays/methods is NOT recommended.

Program Information

- Three 250.0-µL synthetic whole blood specimens
- Two shipments per year



Red Blood Cell Antigen Genotyping RAG				
Procedure	Program Code	Challenges per Shipment		
	RAG			
RBC blood group genotyping for phenotype prediction	ı	3		

Program Information

- Three 2.0-mL whole blood specimens
- Two shipments per year

Noninvasive Prenatal Testing NIPT **Analyte Program Code** Challenges per Shipment NIPT Cell-free DNA screening for fetal 3 aneuploidy

Noninvasive prenatal testing is an exercise and is not considered proficiency testing. This exercise may be used to meet the requirements for alternative assessment.

- · Three liquid specimens
- · Two shipments per year

Next-Generation Sequencing

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

All laboratories subject to US Clinical Laboratory Improvement Amendments (CLIA) Regulations: Proficiency testing (PT) challenges must NOT be referred to another laboratory for any portion of NGS testing, even if this is how patient testing is routinely performed. For PT challenges, any referral is strictly prohibited by CMS.

Next-Generation Sequencing—Germline NGS				
Procedure	Program Code Challenges per Shipment			
	NGS			
Next-generation sequencing	I	2		

Laboratories will have the ability to analyze more than 100 preselected chromosomal intervals in hg19 (GRCh37) and hg38 (GRCh38) coordinates within various genes; for a full list of genes in this program, please go to cap.org. Under the Laboratory Improvement tab, click on Catalog and Ordering Information. The list is located under the PT Order Supplements header.

Program Information

- One 10.0-µg extracted gDNA specimen; one educational variant interpretation image/ dry challenge
- Methods-based challenge for germline variants for laboratories using gene panels, exome, and genome sequencing
- Two shipments per year

Next-Generation Sequencing—Solid Tumor NGSST				
Procedure	Program Code Challenges per Shipment			
	NGSST			
Next-generation sequencing	I	3		

Additional Information

- This is a methods-based proficiency challenge for laboratories performing targeted next-generation sequencing of cancer genes or mutation hotspots in solid tumors.
- This program includes variants present with a variant allele fraction (VAF) potentially as low as 5%.
- · Paired normal specimen provided

Program Information

- Three 1.0-μg gDNA (50 ng/μL) specimens
- One 3.0-μg gDNA (50 ng/μl) paired normal specimen
- · Two shipments per year

Next-Generation Sequencing—Hematologic Malignancies NGSHM					
Procedure	Program Code Challenges per Shipment				
	NGSHM				
Next-generation sequencing	generation sequencing				

Additional Information

- This is a methods-based proficiency challenge for laboratories performing targeted next-generation sequencing of genes or mutation hotspots in hematologic malignancies.
- This program includes variants present with a variant allele fraction (VAF) potentially as low as 5%.

Program Information

- Three 1.0-μg gDNA (50 ng/μL) specimens
- Two shipments per year

19

Next-Generation Sequencing Solid Tumor Bioinformatics NGSB1 Procedure **Program Code** Challenges per Shipment NGSB1 Illumina TruSight Tumor 15 Panel 1 Illumina TruSight Tumor 170 Panel 1 Illumina TruSight Oncology 500 Panel 1 Thermo Fisher Ion AmpliSeq Cancer 1 Hotspot Panel v2 Thermo Fisher Oncomine 1 ı Comprehensive Assay v3 Thermo Fisher Oncomine 1 Focus Cancer Panel

Additional Information

- This *in silico* bioinformatics program is designed to complement and augment somatic variant wet bench NGS proficiency testing programs by testing a greater diversity of variants at a greater range of variant allele fractions (VAF).
- The BAM and/or FASTQ files are platform-specific and may not be compatible with other instruments/software.
- This program includes variants present with a VAF potentially as low as 5%.
- For platform-agnostic solid tumor bioinformatic proficiency testing challenges, refer to the NGSB4 program, page 268.

- Sequencing files containing somatic variants to be downloaded and incorporated into your laboratory bioinformatics pipeline for analysis and reporting; file sizes range from 100MB to 1GB
- BAM and FASTQ file formats
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Next-Generation Sequencing Solid Tumor Bioinformatics Hybrid NGSB4			
Analyte/Procedure	Program Code Challenges per Shipmen		
	NGSB4		
In silico mutagenized sequencing file(s) containing somatic variants of relevance in solid tumors - platform-agnostic	ı	1	

This is a platform-agnostic hybrid *in silico* proficiency testing and validated materials program for laboratories performing targeted NGS of cancer genes or mutational hotspots in solid tumors.

For panel-specific solid tumor bioinformatic proficiency testing challenges, refer to the NGSB1 program, page 267.

Minimum Requirements:

- Laboratories must provide a gene panel sequencing data file (FASTQ or <u>unaligned</u> BAM) that has been generated using their current clinical sequencing protocols from one of the following sources: A specimen from the NGS Germline program (see page 266) or from one of the following NIST Reference Material cell lines: RM 8398 (NA12878), RM 8391, RM 8392, or RM 8393. FASTQs or <u>unaligned</u> BAMs must be submitted along with a BED file describing the regions targeted and interrogated by your laboratory. Specimens from the NGSST and NGSHM programs or additional Coriell/NIST Reference Material cell lines cannot be used for this program.
- Laboratories can transfer files from most modern browsers/operating systems.
 Due to the extremely large file sizes, 40 Mbps transfer speed or higher is needed to ensure successful transfer of your laboratory's sequencing files to the CAP. For the most up-to-date information on system requirements, click Browser and Operating System Requirements located at the bottom of the cap.org homepage.

Additional Information, Proficiency Testing Program:

 Laboratories will be asked to identify somatic single nucleotide variants and small (1-15bp) insertions, deletions, duplications, and deletions-insertions (delins) in a subset of solid tumor mutational hotspots/genes with VAF potentially as low as 5%. Laboratories will be required to submit results of the variants identified.

Additional Information, Validated Materials:

- The sequencing file will contain up to 75 custom somatic variants that are tailored to the specific assay submitted (depending on the size of the panel provided) at VAF from 3% to 99% (higher allele fractions to mimic loss of heterozygosity or homozygosity) and will include:
 - o Single nucleotide variants
 - o Insertions, deletions, delins, and/or duplications ranging from 1-100bp (1-15bp, 16-50bp, 51-100bp)
 - o For laboratories doing microsatellite instability, microsatellite instability at mono nucleotide tracts in the submitted capture design will be included.

All variants will be modeled based on actual somatic mutations from the COSMIC database. This portion of the program is not traditional proficiency testing and no results will be returned to the CAP; information regarding the variants introduced will be sent along with the mutagenized file.

- · The proficiency testing portion of this program is designed to complement and augment NGS somatic variant wet bench proficiency testing programs by testing for a greater diversity of variants with a wide range of variant allele fractions (VAF), while the validated materials portion is designed to optimize bioinformatics pipelines, augment validations, and assist with pipeline verification after changes to NGS/ bioinformatics processes.
- One panel sequencing data file (FASTQ or <u>unaligned</u> BAM), originating from your laboratory and provided to the CAP, for in silico mutagenesis
- Sequencing files containing somatic variants to be downloaded and analyzed by your laboratory bioinformatics pipeline
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Next-Generation Sequencing Hematologic Malignancies Bioinformatics NGSB3

Procedure	Program Code	Challenges per Shipment
	NGSB3	
Illumina TruSight Myeloid Sequencing Panel	ı	1
Thermo Fisher Oncomine Myeloid Assay		1

Additional Information

- This *in silico* bioinformatics program is designed to complement and augment somatic variant wet bench NGS proficiency testing programs by testing a greater diversity of variants at a greater range of variant allele fractions (VAF).
- The BAM and/or FASTQ files are platform-specific and may not be compatible with other instruments/software.
- This program includes variants present with a VAF potentially as low as 5%.
- For platform-agnostic hematologic malignancies bioinformatic proficiency testing challenges, refer to the NGSB5 program, page 270.

- Sequencing files containing somatic variants to be downloaded and incorporated into your laboratory bioinformatics pipeline for analysis and reporting; file sizes range from 100MB to 1GB
- BAM and FASTQ file formats
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Next-Generation Sequencing Hematologic Malignancies Bioinformatics Hybrid NGSB5

Analyte/Procedure	Program Code	Challenges per Shipment
	NGSB5	
In silico mutagenized sequencing file(s) containing somatic variants of relevance in hematologic malignancies - platform-agnostic	ı	1

This is a platform-agnostic hybrid *in silico* proficiency testing and validated materials program for laboratories performing targeted NGS of cancer genes or mutational hotspots in hematologic malignancies.

For panel-specific hematologic malignancies bioinformatic proficiency testing challenges, refer to the NGSB3 program, page 269.

Minimum Requirements:

- Laboratories must provide a gene panel sequencing data file (FASTQ or <u>unaligned</u> BAM) that has been generated using their current clinical sequencing protocols from one of the following sources: a specimen from the NGS Germline program (see page 266) or from one of the following NIST Reference Material cell lines: RM 8398 (NA12878), RM 8391, RM 8392, or RM 8393. FASTQs or <u>unaligned</u> BAMs must be submitted along with a BED file describing the regions targeted and interrogated by your laboratory. Specimens from the NGSST and NGSHM programs or additional Coriell/NIST Reference Material cell lines cannot be used for this program.
- Laboratories can transfer files from most modern browsers/operating systems.
 Due to the extremely large file sizes, 40 Mbps transfer speed or higher is needed to ensure successful transfer of your laboratory's sequencing files to the CAP. For the most up-to-date information on system requirements, click Browser and Operating System Requirements located at the bottom of the cap.org homepage.

Additional Information, Proficiency Testing Program:

 Laboratories will be asked to identify somatic single nucleotide variants and small (1-15bp) insertions, deletions, duplications, and deletions-insertions (delins) in a subset of hematologic malignancies mutational hotspots/genes with VAF potentially as low as 5%. Laboratories will be required to submit results of the variants identified.

Additional Information, Validated Materials:

- The sequencing file will contain up to 75 custom somatic variants that are tailored to the specific assay submitted (depending on the size of the panel provided) at VAF from 3% to 99% (higher allele fractions to mimic loss of heterozygosity or homozygosity) and will include:
 - o Single nucleotide variants
 - o Insertions, deletions, delins, and/or duplications ranging from 1–100bp (1–15bp, 16–50bp, 51–100bp)

All variants will be modeled based on actual somatic mutations from the COSMIC database. This portion of the program is not traditional proficiency testing and no results will be returned to the CAP; information regarding the variants introduced will be sent along with the mutagenized file.

- · The proficiency testing portion of this program is designed to complement and augment NGS somatic variant wet bench proficiency testing programs by testing for a greater diversity of variants with a wide range of variant allele fractions (VAF) while the validated materials portion is designed to optimize bioinformatics pipelines, augment validations, and assist with pipeline verification after changes to NGS/ bioinformatics processes.
- One panel sequencing data file (FASTQ or <u>unaligned</u> BAM), originating from your laboratory and provided to the CAP, for in silico mutagenesis
- Sequencing files containing somatic variants to be downloaded and analyzed by your laboratory bioinformatics pipeline
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Next-Generation Sequencing Undiagnosed Disorders—Exome NGSE						
Analyte/Procedure Program Code Challenges per Shipmen						
	NGSE					
Exome analysis for germline undiagnosed disorders		1				

Additional Information/Minimum Requirements

- This in silico based program will assess the ability of the laboratory to identify
 germline variants responsible for a provided clinic phenotype as is encountered in
 an undiagnosed disease scenario. In addition to analyzing the in silico mutagenized
 file to identify a genetic diagnosis for the provided clinical scenario, pathogenic or
 likely pathogenic ACMG secondary findings may also be reported.
- Laboratories must provide an exome sequencing data file (FASTQ or <u>unaligned</u> BAM) that has been generated using their current clinical sequencing protocols from one of the following sources: A specimen from the NGS Germline program (see page 266) or from one of the NIST Reference Material cell lines: RM 8398 (NA12878), RM 8391, RM 8392, or RM 8393. Specimens from the NGSST and NGSHM programs or additional Coriell/NIST Reference Material cell lines cannot be used for this program.
- FASTQs or <u>unaligned</u> BAMs must be submitted along with a BED file describing the
 regions targeted and interrogated by your laboratory. Additionally, more than 90%
 of exons targeted and interrogated by your laboratory must have a minimum read
 coverage of 10X.
- Laboratories can transfer files from most modern browsers/operating systems.
 Due to the extremely large file sizes, 40 Mbps transfer speed or higher is needed to ensure successful transfer of your laboratory's sequencing files to the CAP. For the most up-to-date information on system requirements, click Browser and Operating System Requirements located at the bottom of the cap.org homepage.

- One exome sequencing data file, originating from your laboratory and provided to the CAP, for in silico mutagenesis; the mutagenized exome sequencing data file is to be downloaded and analyzed by your bioinformatics pipeline
- The mutagenized exome sequencing file will be accompanied by a clinical history, relevant laboratory data, and results of ancillary studies, where appropriate.
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Next-Generation Sequencing Undiagnosed Disorders—Trio Analysis NGSET

Analyte/Procedure	Program Code	Challenges per Shipment
	NGSET	
Trio (parents and proband) exome analysis for germline undiagnosed disorders	1	3

Additional Information/Minimum Requirements

- This in silico based program will assess the ability of the laboratory to identify germline variants responsible for a provided clinic phenotype in a proband as is encountered in an undiagnosed disease scenario using a trio approach (ie, laboratories will analyze the proband and parents in an effort to determine the diagnosis in the proband). In addition to analyzing the in silico mutagenized files to identify a genetic diagnosis for the provided clinical scenario, inheritance patterns as well as pathogenic or likely pathogenic ACMG secondary findings may also be reported.
- Laboratories must provide exome sequencing data files (FASTQs or <u>unaligned</u> BAMs) that have been generated using their current clinical sequencing protocols from one of the following Genome in a Bottle Consortium trio sources: The Ashkenazi Jewish trio (Coriell IDs GM24385, GM24149, and GM24143 or NIST RM8392) or the Han Chinese trio (Coriell IDs GM24631, GM24694, and GM24695). All exome files must be from the same trio (Ashkenazi Jewish or Han Chinese). Specimens from the NGS, NGSST, and NGSHM programs or additional Coriell/Genome in a Bottle Consortium sources cannot be used for this program.
- FASTQs or <u>unaligned</u> BAMs must be submitted along with a BED file describing
 the regions targeted and interrogated by your laboratory. Additionally, more than
 90% of exons targeted and interrogated by your laboratory must have a minimum
 read coverage of 10X.
- Laboratories can transfer files from most modern browsers/operating systems.
 Due to the extremely large file sizes, 40 Mbps transfer speed or higher is needed to ensure successful transfer of your laboratory's sequencing files to the CAP. For the most up-to-date information on system requirements, click Browser and Operating System Requirements located at the bottom of the cap.org homepage.

- Three exome sequencing data files (one from each parent plus the proband), originating from your laboratory and provided to the CAP, for in silico mutagenesis; the mutagenized exome sequencing data files are to be downloaded and analyzed by your bioinformatics pipeline
- The mutagenized exome sequencing files will be accompanied by a clinical history, relevant laboratory data, and results of ancillary studies, where appropriate.
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Copy Number Variant—Solid Tumor CNVST			
Procedure Program Code Challenges per Shipme			
	CNVST		
Copy number variant—solid tumor	ı	3	

- This program is designed for laboratories using next-generation sequencing for copy number analysis.
- Laboratories will be asked to identify copy number alterations in some of these genes: CDKN2A, CDKN2B, EGFR, ERBB2, FGFR3, MET, MYC, MYCN, TP53.
- Copy number alterations tested will include amplification, gain, copy neutral loss of heterozygosity, and deletion.

Tumor Mutational Burden TMB					
Procedure Program Code Challenges per Shipmer					
ТМВ					
Tumor mutational burden 3					

Additional Information

- This program is intended for laboratories using next-generation sequencing to determine tumor mutational burden.
- This program is appropriate for laboratories using targeted panels and whole exome sequencing.
- Paired normal tissue is included.
- · Specimens are 50% tumor.

Program Information

- One 20-μL gDNA (10ng/μL) specimen
- Two snap-frozen cell pellets
- Two shipments per year

- Three 10-µL gDNA (50ng/µL) specimens
- Three 10-µL gDNA (50ng/µL) paired normal tissues
- Two shipments per year

Molecular Oncology—Solid Tumors

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Microsatellite Instability MSI			
Procedure Program Code Challenges per S			
	MSI		
Microsatellite instability testing (DNA amplification)	1	3	
MLH1 promoter methylation analysis	3		

Laboratories performing DNA mismatch repair assessment by immunohistochemistry methods should see program MMR on page 299.

Program Information

- Three specimens each containing two 10.0-micron unstained paraffin section slides and one H&E slide
- For laboratories performing molecular testing using PCR and NGS
- Two shipments per year

In Situ Hybridization ISH, ISH2			
Analyte/Procedure	nalyte/Procedure Program Code Challenges per Shipme		
	ISH	ISH2	
Epstein-Barr virus (EBV)			4
Human papillomavirus (HPV)			4
Kappa/Lambda (IGK/IGL)			4
ERBB2 (HER2) gene amplification (brightfield)			10

Laboratories performing FISH for interphase chromosomal targets in paraffin sections refer to the Cytogenetics programs, page 255.

These programs are only for laboratories that perform both hybridization and interpretation under the same CLIA number.

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Pro	oram	Intorm	Iation
	EI GIII		ıacıvı

• ISH -

EBV, HPV: Three 4-core tissue microarray slides and one H&E slide (each)

Kappa/Lambda: Four 4-core tissue microarray slides and one H&E slide

- ISH2 Two 5-core tissue microarray slides in duplicate
- Two shipments per year

DNA Extraction & Amplification FFPE MH05					
Procedure	Program Code Challenges per Shipment				
MH05					
DNA purification		1			

This is a methods-based proficiency challenge to examine DNA purification from formalin-fixed, paraffin-embedded (FFPE) tissues. Laboratories will be able to purify DNA from FFPE sections and amplify control targets using laboratory-provided reagents.

Program Information

- Three 10.0-micron paraffin sections
- · Two shipments per year

19

Neoplastic Cellularity NEO					
Procedure Program Code Challenges per Shipm					
NEO					
Online assessment of percent neoplastic cellularity	ı	10			

Program Information

- Ten regions of interest (ROIs) using online whole slide images
- A method-based preanalytic program to assess competency for determining percent neoplastic cellularity
- Powered by DigitalScope® technology
- Individual reporting fields for up to five pathologists are available.
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available

Sarcoma Fusion Gene SARC				
Gene	Program Code Challenges per Shipment			
	SARC			
Sarcoma fusion gene*		3		

^{*}See fusion gene listing below.

Laboratories performing FISH for sarcoma translocation refer to the Cytogenetics programs, page 255.

Program Information

- Three snap-frozen cell pellets from which approximately 5.0-µg of RNA can be extracted
- For laboratories performing molecular testing using RT-PCR and NanoString
- · Two shipments per year

Sarcoma Fusion Gene Listing

COL1A1::PDGFB, t(17;22)	EWSR1::FLI1 or EWSR1::ERG	PAX3::FOXO1 or PAX7::FOXO1
ETV6::NTRK3, t(12;15)	EWSR1::WT1, t(11;22)	SS18::SSX1, t(X;18)
EWSR1::ATF1, t(12;22)	FUS::DDIT3, t(12;16)	SS18::SSX2, t(X;18)
EWSR1::ERG, t(21;22)	PAX3::FOXO1, t(2;13)	SS18::SSX1 or SS18::SSX2
EWSR1::FLI1, t(11;22)	PAX7::FOXO1, t(1;13)	

19

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Cell-Free Tumor DNA CFDNA					
Analyte/Procedure Program Code Challenges per Shipme					
	CFDNA				
cfDNA	I	3			

- · DNA fragments stabilized in simulated plasma
- This is not intended for laboratories that perform circulating tumor cell (CTC) analysis.
- Genes in this program include: EGFR, BRAF, KRAS, NRAS, IDH1, PIK3CA, ERBB2, MET, ALK, and BRCA1.
- This program includes variants present with a VAF range of 0.1% 3.0%.

Program Information

- Three 125-ng DNA (25 ng/mL) specimens
- · Two shipments per year

Fusion RNA Sequencing RNA					
Analyte/Procedure	Program Code	Challenges per Shipment			
RNA		3			

Additional Information

- Total RNA from a cell line engineered to contain desired fusion RNA
- This is for laboratories using RNAseq to detect gene fusion transcripts.
- This is not intended to replace the current program (SARC) for reverse transcription (RT)-PCR based detection (see page 275).
- Potential fusion variants include: CD74::ROS1, EML4::ALK, ETV6::NTRK3, FGFR3::TACC3, PAX8::PPARG, and SLC45A3::BRAF.
- · Specific intragenic fusion/exon skipping variants may also be included, specifically EGFRvIII and MET exon 14 skipping.

Solid Tumor—Other BRAF, EGFR, KRAS, KIT						
Analyte		Program Code Challenges per Shipment				
	BRAF	BRAF EGFR KRAS KIT				
BRAF					3	
EGFR		ı			3	
KRAS	3				3	
KIT					3	
PDGFRA				ı	3	

Program Information

- Three 500-ng RNA (20 ng/μL) specimens
- · Two shipments per year

Program Information

- · BRAF, EGFR, KRAS -Paraffin-embedded sections or shavings
- KIT -

One specimen containing four 10.0-micron unstained paraffin section slides and one H&E slide

Two 1.0-µg gDNA (50 ng/µL) specimens

- For laboratories performing molecular testing using PCR
- · Two shipments per year

Multigene Tumor Panel MTP					
Analyte	Program Code	Challenges per Shipment			
	MTP				
BRAF	1	3			
EGFR	1	3			
ERBB2 (HER2)	I	3			
KIT	1	3			
KRAS	1	3			
NRAS	I	3			
PDGFRA	I	3			
PIK3CA		3			

CAP accredited laboratories that perform testing for the detection of somatic single nucleotide variants, insertions, and deletions in *BRAF*, *EGFR*, and *KRAS* by non-NGS methods are required to enroll in either MTP or the respective single gene programs. This includes laboratories that perform non-NGS-based multiplexed assays and nonmultiplexed assays (eg, Sanger sequencing). Laboratories that perform NGS-based testing of somatic single nucleotide variants, insertions, and deletions in *BRAF*, *KRAS*, *EGFR*, and/or other genes are required to enroll in NGSST (on page 266) as this proficiency testing program provides challenges with lower variant allele fractions (VAF) as well as challenges in other genes commonly included in NGS-based panels for the identification of somatic variants in solid tumors.

Glioma GLI					
Analyte Program Code Challenges per Shi					
GLI					
MGMT		3			
IDH1, IDH2	I	3			

Program Information

- Three 2.0-µg gDNA
 (50 ng/µL) specimens for
 laboratories performing
 molecular testing on multiple
 targets
- · Two shipments per year

- Four 2.0-μg gDNA (50 ng/μL) specimens
- One specimen containing four 10.0-micron unstained paraffin section slides and one H&E slide
- For laboratories performing molecular testing using PCR
- · Two shipments per year

Molecular Oncology—Hematologic

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Molecular Hematologic Oncology MHO/MHO1, MHO2/MHO3, MHO5					
Procedure/Gene		Program Code		Challenges per Shipment	
	MHO/MHO1	MH02/MH03	MH05		
Lymphoid Malignancy Genotyp	ing				
IGH				3	
IGH::BCL2 major				3	
IGH::BCL2 minor				3	
IGH::CCND1				3	
IGK				3	
TRB				3	
TRG				3	
Myeloid Malignancy Genotypin	g				
BCR::ABL1 p190				3	
BCR::ABL1 p210		•		3	
CALR				3	
CBFB::MYH11				3	
FLT3 ITD		•		3	
FLT3 TKD				3	
JAK2 c.1849G>T p.V617F				3	
KMT2A-PTD (MLL-PTD)				3	
MPL				3	
NPM1				3	
PML::RARA		•		3	
RUNX1::RUNX1T1		I		3	
DNA extraction and amplification from formalin-fixed, paraffin-embedded (FFPE) tissue			ı	1	

- MHO One sample vial containing purified DNA (200 µg/mL per vial) for each specimen
- MH01 MH0 specimens in duplicate for additional DNA testing
- MHO2 Two sample vials; one with purified DNA containing 200 µg/mL and one with purified RNA containing 400 µg/mL
- MH03 MH02 specimen in duplicate for additional DNA and RNA testing
- MH05 Three 10.0-micron paraffin sections; extraction and amplification from FFPE tissue will be assessed by a method-based challenge
- Two shipments per year; ships on dry ice (dry ice does not apply to MHO5)

IGHV Mutation Analysis IGHV			
Analyte/Procedure	Program Code	Challenges per Shipment	
IGHV		3	

- Sequence analysis of the clonal immunoglobulin heavy chain V gene (IGHV) to determine somatic hypermutation (SHM) status
- Any sequencing method may be used.
- Report productive/unproductive rearrangement, SHM status, percent similarity, and V-gene utilization.

Measurable (Minimal) Residual Disease MRD, MRD1, MRD2				
Analyte		Program Cod	Challenges per Shipment	
	MRD	MRD1	MRD2	
BCR::ABL1 p190				3
BCR::ABL1 p210				3
PML::RARA			ı	3

Program Information

- Three 20-μg DNA specimens (200 ng/μL)
- Two shipments per year

- MRD, MRD1, MRD2 Three RNA specimens in sterile water
- For laboratories diagnosing and monitoring leukemia tumor burden by measuring the quantity of BCR::ABL1 or PML::RARA fusion transcripts
- Two shipments per year; ships on dry ice

Navigating Multimodality Biomarker Assessment NMBA/NMB1				
Program Name	Program Code	Cases per Mailing		
NMBA/NMB1				
Multimodality biomarker assessment case analysis	ı	2		

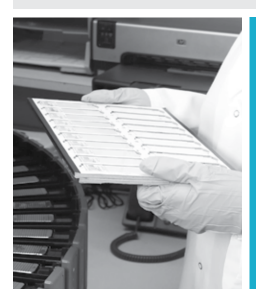
Biomarkers can be tested through a variety of methodologies (eg, flow cytometry, immunohistochemistry, next-generation sequencing, PCR-based testing, karyotype, FISH, clinical chemistry). Each modality has different technical strengths and weaknesses, along with varied analytical and clinical specifications such as sensitivity and specificity. Results may not be clearly concordant across the modalities. This program will challenge pathologists to resolve discrepancies between different methods or "multimodality" biomarker testing.

- · NMBA Online program, whole slide images powered by DigitalScope technology (if available); NMBA provides CME or CE for one pathologist or laboratory professional
- NMB1 Reporting option with CME or CE credit for each additional pathologist or laboratory professional (within the same institution); must order in conjunction with program NMBA
- · Two mailings per year with two cases each mailing
- · Earn a maximum of four CME credits (AMA PRA Category 1 Credits™) per pathologist and a maximum of four CE credits per laboratory professional per year.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Anatomic Pathology

20 Anatomic Pathology



Prepare for success with our PAP PT and PAP Education programs.

- Every slide is reviewed and approved by pathologists and cytotechnologists before it is put in circulation.
- All slide sets are reviewed every six months by a staff cytotechnologist.
- Slides that do not maintain consensus grading are removed from the program and reviewed by a committee of pathologist experts.

Anatomic Pathology

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Surgical Pathology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Online Performance Improvement Program in Surgical Pathology PIPW/PIPW1			
Program	Program Code Challenges per Shipment		
	PIPW/PIPW1		
Surgical pathology case review		10	

Additional Information

- Program PIPW prepares pathologists to succeed by providing ongoing diagnostic learning in general surgical pathology.
- Pathologists can assess their diagnostic skills and compare their performance with that of their peers.
- Included PIPW case selections feature:
 - O A variety of neoplastic and nonneoplastic lesions
 - o Inflammatory and infectious diseases
 - O Various sites, encompassing a variety of organ systems
 - Two PIPW cases per release are from smaller tumors and do not duplicate PIP (glass).
- See system requirements on page 12.

- PIPW Ten diagnostic challenges/whole slide H&E images with clinical history; CME credit is available for one pathologist; for each additional pathologist, order PIPW1
- PIPW1 Reporting option with CME credit for each additional pathologist (within the same institution); must order in conjunction with program PIPW
- Earn a maximum of 40 CME credits (AMA PRA Category 1 Credits™) per pathologist for completion of an entire year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Powered by DigitalScope® technology
- Four online activities per year; your CAP shipping contact will be notified via email when the activity is available



Performance Improvement Program in Surgical Pathology PIP/PIP1				
Program Code Challenges per Shipment				
PIP/PIP1				
Surgical pathology case review ■ 10				

- PIP prepares pathologists to succeed by providing ongoing diagnostic learning in general surgical pathology.
- · This program:
 - o Provides a practical approach to continuing education
 - o Gives pathologists a method to assess their diagnostic skills and compare their performance with that of their peers
 - O Allows staff to experience smaller tumors and more interesting cases by providing three online cases per release
 - o Features PIP case selections that include:
 - A variety of neoplastic and nonneoplastic lesions
 - Inflammatory and infectious diseases
 - Various sites, encompassing a variety of organ systems

- PIP Ten diagnostic challenges with clinical history: seven H&E stained glass slides and three online only cases; CME credit is available for one pathologist; for each additional pathologist, order PIP1
- · PIP1 Reporting option with CME credit for each additional pathologist (within the same institution); must order in conjunction with program PIP
- · Powered by DigitalScope technology
- · Earn a maximum of 40 CME credits (AMA PRA Category 1 Credits) per pathologist for completion of an entire year.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Four shipments per year



Virtual Biopsy Program VBP/VBP1				
Program	Program Code Challenges per Shipment			
	VBP/VBP1			
Online biopsy case review		5		

- VBP prepares pathologists to succeed by providing ongoing diagnostic learning in surgical pathology.
- This program is applicable to all pathologists, including general pathologists, and focuses on biopsy material. Cases may include gross, radiographic, or endoscopic images.
- There are four topical releases per year that focus on benign and malignant pathology. Cases are from selected organ systems and may include a variety of specimen types (eg, core biopsies, endoscopic biopsies, curettings, aspirate smears)
- See system requirements on page 12.

- VBP Five diagnostic challenges/whole slide images with clinical history; reporting with CME credit is available for one pathologist; for each additional pathologist, order VBP1
- VBP1 Reporting option with CME credit for each additional pathologist (within the same institution); must order in conjunction with program VBP
- Earn a maximum of 25 CME credits (AMA PRA Category 1 Credits) per pathologist for completion of an entire year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Powered by DigitalScope technology
- Four online activities per year; your CAP shipping contact will be notified via email when the activity is available



Pathologists can keep abreast of current scientific knowledge with interactive, case-based learning addressing common issues faced in the laboratory.

CPIP supports clinical pathologists as well as anatomic pathologists who cover clinical pathology. A diverse portfolio of real-life case scenarios, including images and clinical background, helps pathologists to stay current on issues and advances in the laboratory.

CPIP is designed for pathologists, by pathologists. Each case is developed and peer-reviewed, ensuring learning is practical and easily applied to work. Thought-provoking questions with feedback and multiple choice knowledge checks assess and confirm diagnostic skills. Participants may apply 1.25 CME credits for each CPIP toward the ABPath's Continuing Certification (CC) requirements.

Clinical Pathology Improvement Program CPIP/CPIP1		
Program Name	Program Code	Cases per Year
	CPIP/CPIP1	
Online cases in clinical pathology	ı	12

Consider CPIP for:

- Medical directors seeking to continuously improve the collective skills and clinical pathology knowledge of their team
- · Pathologists with clinical and/or laboratory management responsibilities
- Pathologists seeking CME CC credits in clinical pathology
- · Subspecialty clinical pathologists who need to keep current

Discipline	Case Schedule (subject to change)	Month 2025
Laboratory Management	Provider Performed Microscopy (PPM) Testing	January
Transfusion Medicine	ABO Discrepancies	February
Transfusion Medicine	Blood Bank Regulations	March
Chemistry	Hyperbilirubinemia	April
Hematology	Hematologic Pleural Effusions	May
Microbiology	Bloodstream Infections	June
Laboratory Management	CLIA Director Responsibilities and Risks	July
Cytogenetics	Plasma Cell Myeloma	August
Transfusion Medicine	Transfusion Reactions	September
Molecular Pathology	Pharmacogenomics	October
Hematology	Peripheral Blood Smear - Part 1	November
Hematology	Peripheral Blood Smear - Part 2	December

To learn more visit cap.org and search CPIP.

- CPIP One online clinical laboratory case per month
- CPIP1 Additional pathologist (within the same institution) reporting option with CME credit; must order in conjunction with CPIP
- Earn a maximum of 15 CME credits (AMA PRA Category 1 Credits™) per year.
- Twelve cases per year; your CAP shipping contact will be notified via email when the activity is available



Touch Imprint/Crush Preparation TICP/TICP1			
Procedure Program Code Challenges per St			
	TICP/TICP1		
Online slide and image program in rapid assessment case review		4	

- The TICP program gets surgical pathologists, cytopathologists, and cytotechnologists ready to succeed by familiarizing them with the cytomorphologic features of pathologic processes and tumors in touch imprints and crush or scrape preparations. These specimens are prepared either for intraoperative consultation (frozen section) or rapid on-site evaluation (ROSE) of tissue biopsies for adequacy and/or interpretation. Participants will learn to make an immediate adequacy assessment, assign the process to a general category, and triage the specimen to appropriate ancillary studies. Participants will review digital whole slides of the TICP preparations (hematoxylin & eosin, modified Wright-Giemsa, and/or Papanicolaou stains), static images of the preparation and ancillary studies, and clinical history/radiographic findings to reach a diagnosis. Each case has a complete description of entities in the differential diagnosis along with a discussion of the correct interpretation.
- Participants will receive immediate feedback on interpretations, ancillary studies, and case-related adequate assessment.
- The cases will focus on miscellaneous topics.
- · May include rarely captured cases that may not be available on the glass slide
- See system requirements on page 12.

- TICP Four online
 assessment challenges
 with clinical history; TICP
 provides CME or CE credit
 for one pathologist or
 cytotechnologist; for each
 additional pathologist or
 cytotechnologist, order TICP1
- TICP1 Reporting option with CME or CE credit for each additional pathologist/ cytotechnologist (within the same institution); must order in conjunction with program TICP
- Earn a maximum of 10 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 10 CE credits per cytotechnologist for completion of an entire year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Online whole slide images powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



CAP/NSH HistoQIP HQIP				
Stain/Tissue	Program Code	Challenges per Shipme		
	HQIP	Α	В	
H&E - Liver resection or biopsy		1		
H&E - Breast, needle core biopsy		1		
IHC - Hep-Par, liver resection		1		
IHC - GATA3, breast needle core biopsy		1		
Special Stain - Reticulin, liver resection		1		
H&E - Stomach resection with gastrointestinal stromal tumor (GIST) and nonneoplastic stomach mucosa	ı		1	
H&E - Prostate biopsy, containing both adenocarcinoma and nonneoplastic acini			1	
IHC - DOG1, gastrointestinal stromal tumor (GIST) positive for DOG1			1	
IHC - PIN4, prostate biopsy with adenocarcinoma and nonneoplastic glands	•		1	
Special Stain - Congo Red, positive for amyloid	ı		1	

HistoQIP improves histologic slide preparation in anatomic pathology laboratories. In this educational program, an expert panel of pathologists, histotechnologists, and histotechnicians evaluates submitted slides for histologic technique using uniform grading criteria. Participants receive laboratory-specific evaluations and participant summaries with performance benchmarking, commentary from evaluators, and peer comparison data.

Program Information

- · Participant laboratories may submit up to five stained coverslipped glass slides (one from each category) per mailing.
- · Includes photographs
- · Two shipments per year



CAP/NSH Pediatric Prog	HistoQIP gram HQPED		NEW
Stain/Tissue	Program Code	Challenges p	oer Shipment
	HQPED	Α	В
H&E - Colon resection for Hirschsprung disease	ı	1	
IHC - Calretinin, colon resection for Hirschsprung disease	ı	1	
H&E - Wilms tumor, renal resection	1	1	
IHC - WT1, Wilms tumor, renal resection	ı	1	
H&E - Rhabdomyosarcoma	I		1
IHC - Myogenin, positive rhabdomyosarcoma	ı		1
H&E - Hepatoblastoma, liver resection	ı		1
IHC - B-catenin, hepatoblastoma liver resection	ı		1

Program Information

- Participant laboratories may submit up to four stained coverslipped glass slides (one from each category) per mailing.
- Two shipments per year



CAP/NSH HistoQIP Cell Block Preparations HQCLB					
Stain/Tissue	Program Code	Challenges per Shipme			
	HQCLB	Α	В		
H&E - Neck mass, HPV on squamous cell carcinoma		1			
IHC - p16, squamous cell carcinoma		1			
H&E - Pleural fluid, metastatic breast ductal carcinoma		1			
IHC - GATA3, metastatic breast ductal carcinoma	•	1			
H&E - Lung mass, FNA on a lung squamous cell carcinoma	•		1		
IHC - p40 or p63, lung mass, FNA on a lung squamous cell carcinoma			1		
H&E - Peritoneal fluid, positive PAX8 metastatic carcinoma	•		1		
IHC - PAX8, metastatic carcinoma			1		

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology and cytopathology laboratories that handle cell block preparations.

Program Information

- Participants may submit up to four stained coverslipped glass slides (one from each category) per mailing.
- Two shipments per year



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CAP/NSH HistoQIP Targeted Therapy HQTAR					
Stain/Tissue	Program Code	Challenges p	er Shipment		
	HQTAR	Α	В		
H&E - Breast ductal carcinoma, core needle biopsy	ı	1			
IHC - HER2, breast ductal carcinoma, core needle biopsy	1	1			
H&E - Breast resection, lobular carcinoma	I	1			
IHC - ER, breast resection, lobular carcinoma	I	1			
H&E - Gastroesophageal adenocarcinoma	I		1		
IHC - HER2, gastroesophageal adenocarcinoma	I		1		
H&E - Gastroesophageal adenocarcinoma	I		1		
IHC - Claudin 18.2, gastroesophageal adenocarcinoma	ı		1		

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology laboratories that handle specimens undergoing analysis for targeted therapies.

Program Information

- · Participant laboratories may submit up to four stained coverslipped glass slides (one from each category) per mailing.
- · Two shipments per year



CAP/NSH HistoQIP Whole Slide Image Quality Improvement Program HQWSI Stain/Tissue **Program Code** Challenges per Shipment HQWSI В Α H&E - Kidney biopsy H&E - Pancreas resection 1 IHC - Synaptophysin, pancreas resection 1 Special Stain - Silver (Jones), kidney biopsy 1 H&E - Prostate, invasive adenocarcinoma 1 biopsy H&E - Ovary resection 1 H&E - Lung biopsy 1 IHC - TTF-1, lung biopsy 1 Special Stain - AFB, control tissue 1 H&E - Breast, invasive carcinoma, resection 1 or biopsy

The program provides feedback to laboratories using whole slide imaging for clinical applications. Participants upload their scanned whole slide images to the CAP designated server. In this educational program, an expert panel of pathologists, histotechnologists, and histotechnicians evaluates whole slide images for histologic technique and image quality. Participants receive laboratoryspecific evaluations and participant summaries with performance benchmarking, commentary from evaluators, and peer comparison data as well as annotated feedback directly on their uploaded images.

Program Information

- Participant laboratories may submit up to five stained coverslipped glass slides and corresponding scanned whole slide images per mailing.
- Online whole slide images powered by DigitalScope technology
- · Two shipments per year



CAP/NSH HistoQIP Biopsy Series HQIPBX				
Stain/Tissue	Program Code	Challenges p	er Shipment	
	HQIPBX	Α	В	
H&E - Bladder biopsy	I	1		
H&E - Cervical biopsy	1	1		
H&E - Skin punch biopsy	1	1		
H&E - Stomach biopsy	1	1		
H&E - Colon biopsy	1		1	
H&E - Endometrial biopsy	1		1	
H&E - Prostate needle biopsy	I		1	
H&E - Breast core biopsy	ı		1	

The HistoQIP Biopsy Series is an additional program to improve the preparation of histologic slides in all anatomic pathology laboratories. In this educational program, an expert panel of pathologists, histotechnologists, and histotechnicians evaluates submitted slides for histologic technique using uniform grading criteria. Participants receive laboratory-specific evaluations and participant summaries with performance benchmarking, commentary from evaluators, and peer comparison data.

Program Information

- Participants may submit up to four H&E stained and coverslipped glass slides (one from each category) per mailing.
- Two shipments per year



CAP/NSH HistoQIP Specialty Series HQBX1, HQBX2, HQBX3, HQBX4						
Stain/Tissue		Progra	m Code		Challenges per Shipment	
	HQBX1	HQBX2	HQBX3	HQBX4	A	В
Gastrointestinal Biopsy Module						
H&E - Colon biopsy					1	1
H&E - Esophagus biopsy					1	1
H&E - Small intestine biopsy					1	1
H&E - Stomach biopsy	•				1	1
Dermatologic Biopsy Module						
H&E - Alopecia biopsy					1	1
H&E - Skin excisional biopsy (large excision)		•			1	1
H&E - Skin punch biopsy					1	1
H&E - Skin shave biopsy					1	1
Urogenital Tract Biopsy Module						
H&E - Bladder biopsy (nonneoplastic)			•		1	1
H&E - Bladder biopsy (with urothelial carcinoma)					1	1
H&E - Prostate needle biopsy (nonneoplastic)					1	1
H&E - Prostate needle biopsy (with carcinoma)					1	1
Gynecological Biopsy Module						
H&E - Cervical biopsy					1	1
H&E - Endometrial biopsy					1	1
H&E - Cervical cone/LEEP					1	1
H&E - Vulvar biopsy					1	1

The HistoQIP Specialty Series includes modules to improve the preparation of histologic slides in all anatomic pathology laboratories that handle gastrointestinal, dermatologic, urogenital tract, and gynecologic biopsies. In this educational program, an expert panel of pathologists, histotechnologists, and histotechnicians evaluates submitted slides for histologic technique using uniform grading criteria. Participants receive laboratory-specific evaluations and participant summaries with performance benchmarking, commentary from evaluators, and peer comparison data.

Program Information

- HQBX1, HQBX2, HQBX3, HQBX4 - Participants may submit up to four H&E stained and coverslipped glass slides (one from each category) per mailing
- Two shipments per year



CAP/NSH HistoQIP In Situ Hybridization (Kappa/Lambda) HQISH				
Stain/Tissue	Program Code	Challenges p	er Shipment	
	HQISH	Α	В	
H&E - Nonneoplastic lymph node excision (not a biopsy)		1		
ISH - DNA/RNA negative control probe ISH	1	1		
ISH - DNA/RNA positive control probe ISH	1	1		
ISH - Kappa ISH (Kappa probe, ISH)	•	1		
ISH - Lambda ISH (Lambda probe, ISH)	1	1		
H&E - Bone marrow core biopsy	•		1	
ISH - DNA/RNA negative control probe ISH	•		1	
ISH - DNA/RNA positive control probe ISH	1		1	
ISH - Kappa ISH (Kappa probe, ISH)			1	
ISH - Lambda ISH (Lambda probe, ISH)			1	

This program augments efforts to improve the preparation of ISH slides in all anatomic pathology laboratories that handle specimens undergoing analysis for Kappa and Lambda detection by chromogenic in situ hybridization.

Program Information

- Participant laboratories may submit up to four stained coverslipped glass slides (one from each category) per mailing.
- · Two shipments per year



CAP/NSH HistoQIP IHC Series HQIHC					
Stain/Tissue	Program Code	Challenges p	er Shipment		
	HQIHC	Α	В		
IHC - CD138, plasmacytoma	I	1			
IHC - CD3, nonneoplastic colonic biopsy		1			
IHC - CK5/6, skin biopsy		1			
IHC - INSM1, small cell carcinoma	I	1			
IHC - SAT6, solitary fibrous tumor		1			
IHC - CD23, lymph node excision	I		1		
IHC - CD34, skin biopsy	I		1		
IHC - p53, ovarian serous carcinoma			1		
IHC - TRPS1, breast carcinoma			1		
IHC - Myogenin, skeletal muscle			1		

The HistoQIP IHC series improves the preparation of immunohistochemistry slides in all anatomic laboratories that handle a broad range of surgical specimens. In this educational program, an expert panel of pathologists, histotechnologists, and histotechnicians evaluates submitted slides for histologic technique using uniform grading criteria. Participants receive laboratory-specific evaluations and participant summaries with performance benchmarking, commentary from evaluators, and peer comparison data.

Program Information

- Participants may submit up to five stained coverslipped slides (one from each category) per mailing.
- · Two shipments per year



CAP/NSH HistoQIP Central Nervous System IHC HQNEU Stain/Tissue Challenges per Shipment **Program Code** В **HQNEU** Α H&E - Meningioma 1 IHC - Epithelial membrane antigen (EMA), 1 positive meningioma IHC - Ki-67, positive meningioma 1 H&E - Schwannoma 1 1 IHC - S100, schwannoma ı 1 H&E - IDH1 mutant astrocytoma 1 IHC - ATRX, IDH1 mutant astrocytoma IHC - IDH1, IDH1 mutant astrocytoma 1 H&E - Glioblastoma, IDH-wildtype П 1 IHC - p53, glioblastoma, IDH-wildtype 1

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology laboratories that handle central nervous system gliomas.

Program Information

- Participant laboratories may submit up to five stained coverslipped glass slides (one from each category) per mailing.
- · Two shipments per year



CAP/NSH HistoQIP Non-small Cell Lung Carcinoma IHC HQNSC

Stain/Tissue	Program Code	Challenges per Shipmen	
	HQNSC	A	В
H&E - Lung adenocarcinoma		1	
IHC - TTF-1, lung adenocarcinoma		1	
IHC - Napsin A, lung adenocarcinoma		1	
H&E - ALK, positive lung adenocarcinoma		1	
IHC - ALK, positive lung adenocarcinoma		1	
H&E - Lung squamous cell carcinoma			1
IHC - p40 or p63, lung squamous cell carcinoma	•		1
IHC - CK5 or CK5/6, lung squamous cell carcinoma	•		1
H&E - PD-L1, positive lung squamous cell carcinoma			1
IHC - PD-L1, positive lung squamous cell carcinoma			1

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology laboratories that handle non-small cell lung carcinoma.

Program Information

- Participants may submit up to three IHC and two H&E stained coverslipped glass slides (one from each category) per mailing.
- · Two shipments per year



CAP/NSH HistoQIP Melanoma IHC HQMEL					
Stain/Tissue	Program Code	Challenges p	er Shipment		
	HQMEL	Α	В		
H&E - Melanoma skin biopsy	•	1			
IHC - Melan A/MART-1 melanoma skin biopsy	1	1			
IHC - SOX10, melanoma skin biopsy	•	1			
H&E - PD-L1, positive melanoma skin biopsy	ı	1			
IHC - PD-L1, positive melanoma skin biopsy	1	1			
H&E - Melanoma skin resection	•		1		
IHC - S100, melanoma skin resection	•		1		
IHC - HMB-45, melanoma skin resection	1		1		
H&E - Nevus resection	I		1		
IHC - p16, nevus resection	I		1		

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology laboratories that handle skin specimens containing melanoma.

Program Information

- Participant laboratories may submit up to five stained coverslipped glass slides (one from each category) per mailing.
- Two shipments per year



CAP/NSH HistoQIP Mismatch Repair IHC HQMMR					
Stain/Tissue	Program Code	Code Challenges per Shipme			
	HQMMR	Α	В		
H&E - Colonic adenocarcinoma		1			
IHC - MLH1, colonic adenocarcinoma		1			
IHC - MSH2, colonic adenocarcinoma		1			
IHC - MSH6, colonic adenocarcinoma		1			
IHC - PMS2, colonic adenocarcinoma		1			
H&E - Endometrial adenocarcinoma			1		
IHC - MLH1, endometrial adenocarcinoma			1		
IHC - MSH2, endometrial adenocarcinoma			1		
IHC - MSH6, endometrial adenocarcinoma			1		
IHC - PMS2, endometrial adenocarcinoma			1		

This program augments efforts to improve the preparation of H&E and immunohistochemical slides in all anatomic pathology laboratories that handle colonic and endometrial tumors performing mismatch repair IHC.

Program Information

- Participants may submit up to four IHC and one H&E stained coverslipped glass slides (one from each category) per mailing.
- · Two shipments per year



General Immunohistochemistry

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Immunohistochemistry MK		
Procedure	Program Code	Challenges per Shipment
	MK	
Immunohistochemistry	I	16

The MK program allows laboratories to compare their assay methodology and results with all participating laboratories. Case materials are donated and represent a variety of diagnostic entities. Markers will vary in each case and will provide a wide range of IHC testing for routine surgical pathology practices.

Program Information

- · Five glass slides with unstained tissue sections from four separate cases; each case includes four slides for selected IHC markers and one slide for H&E
- Two shipments per year

CD117 Immunohistochemistry Tissue Microarray PM1			
Analyte Program Code Challenges per Shipment			
	PM1		
CD117	ı	10	

For ER/PgR testing, see the PM2 program on page 297.

ram Code	Challanges nor Chinment
	Challenges per Shipment
PM1	
1	10
_	PM1

Immunohistochemistry Tissue Microarray Series PM5			
Analyte Program Code Challenges per Shipmer			
	PM5		
Folate 1 Receptor	I	10	
SS18	I	10	

Each year, the PM5 program will feature two different markers for immunohistochemistry laboratories to evaluate assay performance on a variety of tissue and/or tumor types. The IHC markers for this program may change from those listed above due to development constraints.

Program Information

- One 10-core tissue microarray slide
- · One shipment per year

Program Information

- Two 10-core tissue microarray slides, one for Folate 1 receptor and one for SS18
- · One shipment per year

These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

p53 Immunohistochemistry Tissue Microarray P53		
Analyte	Program Code	Challenges per Shipment
	P53	
p53		10

The purpose of this program is to assess the laboratory's ability to detect various patterns of p53 staining, which is diagnostically useful in several tumor types.

Program Information

- One 10-core tissue microarray slide
- · Two shipments per year

Dermatopathology Immunohistochemistry DPIHC			
Procedure Program Code Challenges per Shipmer			
	DPIHC		
Dermatopathology		8	

This case-based program assesses the laboratory's ability to perform and interpret immunostains commonly used in dermatopathology practice.

Program Information

- Six glass slides with unstained tissue sections from two separate cases; each case includes four slides for selected IHC markers, one slide for H&E, and one slide for negative control
- Two shipments per year

CAP/ACMG <i>ERBB2 (HER2)</i> Amplification by FISH, Interpretation Only CYHI			
Analyte/Procedure Program Code Challenges per Shipment			
	СҮНІ		
ERBB2 (HER2) amplification in breast cancer, interpretation only		3	

Additional Information

- ERBB2 (HER2) Amplification by FISH, Interpretation Only is not considered proficiency testing. This exercise may be used to meet the requirements for alternative assessment.
- This program is for laboratories that perform <u>interpretation only</u> for *ERBB2 (HER2)* FISH for breast cancer.
- For laboratories that perform both hybridization and interpretation for *ERBB2* (*HER2*) FISH for breast cancer under the same CLIA number, see program CYH on page 255.

Program Information

- Three online interpretation challenges; your CAP shipping contact will be notified <u>via email</u> when the activity is available
- Two shipments per year



These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

Immunohistochemistry Predictive Markers

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

HER2 Immunohistochemistry HER2		
Analyte	Program Code	Challenges per Shipment
	HER2	
HER2		20

The HER2 program fulfills the proficiency testing requirement stated in the ASCO/CAP HER2 Testing Guideline. Due to the unique nature of these human donor-based materials, the shipping date is subject to change. If this should occur, the CAP will provide notification prior to the originally scheduled shipping date.

Program Information

- Two 10-core tissue microarray slides
- · Two shipments per year

Gastric HER2 GHER2		
Analyte	Program Code	Challenges per Shipment
	GHER2	
HER2		10

Additional Information

- The Gastric HER2 program fulfills the proficiency testing requirement stated in the CAP/ASCP/ASCO Gastroesophageal HER2 Testing Guideline.
- The interpretive criteria for HER2 immunohistochemistry performed on gastroesophageal adenocarcinomas differs significantly from breast carcinoma.
 The GHER2 program will help participating laboratories understand these differences.

ER/PgR Immunohistochemistry Tissue Microarray PM2			
Analyte Program Code Challenges per Shipmen			
	PM2		
Estrogen receptor (ER)	I	20	
Progesterone receptor (PgR)	I	20	

The PM2 program fulfills the ER proficiency testing requirement and the PgR alternative assessment requirement stated in the ASCO/CAP ER/PgR Testing Guideline. Due to the unique nature of these human donor-based materials, the shipping date is subject to change. If this should occur, the CAP will provide notification prior to the originally scheduled shipping date.

Program Information

- One 10-core tissue microarray slide
- · Two shipments per year

Program Information

- Four 10-core microarray slides, two for ER and two for PgR
- · Two shipments per year

These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

HER2 and ER Immunohistochemistry Interpretation Only HERI			
Analyte/Procedure Program Code Challenges per Shipment			
	HERI		
HER2 online image review	I	10	
ER online image review	I	10	

- · HER2 and ER Immunohistochemistry Interpretation Only is an exercise and is not considered proficiency testing.
- This program is for laboratories that perform interpretation only for HER2 and ER for breast cancer and may be used for quality assessment.
- For laboratories that perform both staining and interpretation for HER2 and ER for breast cancer under the same CLIA number, see page 297.

• Ten online whole slide images for HER2 by IHC interpretation only

Program Information

- Ten online whole slide images for ER by IHC interpretation only
- · Powered by DigitalScope technology
- Ten whole slide H&E images for HER2 and ER
- · One online activity per year; your CAP shipping contact will be notified via email when the activity is available

CD20 Immunohistochemistry Tissue Microarray PM3		
Analyte	Program Code	Challenges per Shipment
	PM3	
CD20		10

For ER/PgR testing, see the PM2 program on page 297.

Program Information

- One 10-core tissue microarray slide
- · Two shipments per year

Highly Sensitive Anaplastic Lymphoma Kinase IHC PM6			
Analyte	Program Code	Challenges per Shipment	
	PM6		
Highly sensitive anaplastic lymphoma kinase IHC (ALK)	I	10	

This program assesses the laboratory's ability to detect ALK-rearranged lung cancers using highly sensitive ALK immunohistochemistry. The ALK1 clone is NOT highly sensitive and should not be used in this program.

Program Information

- One 10-core tissue microarray slide
- · Two shipments per year

BRAF V600E BRAFV			
Analyte Program Code Challenges per Shipme			
	BRAFV		
BRAF V600E		10	

The purpose of this program is to assess the laboratory's ability to detect BRAF V600E mutant tumors using mutation-specific immunohistochemistry.

Program Information

- One 10-core tissue microarray slide
- · Two shipments per year

These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

CD30 Immunohistochemistry Tissue Microarray CD30				
Analyte	Program Code Challenges per Shipment			
	CD30			
CD30	I	10		

This program assesses the laboratory's ability to detect CD30 expression in lymphomas, which has emerged as a key therapeutic target.

Program InformationOne 10-core tissue

- One 10-core tissue microarray slide
- Two shipments per year

DNA Mismatch Repair MMR			
Procedure	Program Code	Challenges per Shipment	
	MMR		
MLH1 by IHC	I	10	
MSH2 by IHC	I	10	
MSH6 by IHC	I	10	
PMS2 by IHC	I	10	

If your laboratory performs DNA mismatch repair by molecular methods, see the MSI program on page 274.

Program Information

- Four unstained cell line/ tissue microarray slides for the immunohistochemical analysis of DNA mismatch repair proteins MLH1, MSH2, MSH6, and PMS2
- Two shipments per year

PD-L1 Immunohistochemistry PDL1			
Analyte	te Program Code		
	PDL1		
PD-L1	I	10	

The purpose of this program is to assess the laboratory's ability to detect PD-L1 expression and apply various PD-L1 scoring systems.

Program Information

- One 10-core tissue microarray slide; additional slide provided for H&E
- Two shipments per year

These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

Navigating Multimodality Biomarker Assessment NMBA/NMB1			
Program Name		Program Code	Cases per Mailing
		NMBA/NMB1	
Multimodality biomarker assessment case analysis		ı	2

Biomarkers can be tested through a variety of methodologies (eg, flow cytometry, immunohistochemistry, next-generation sequencing, PCR-based testing, karyotype, FISH, clinical chemistry). Each modality has different technical strengths and weaknesses, along with varied analytical and clinical specifications such as sensitivity and specificity. Results may not be clearly concordant across the modalities. This program will challenge pathologists to resolve discrepancies between different methods or "multimodality" biomarker testing.

- NMBA Online program, whole slide images powered by DigitalScope technology (if available); NMBA provides CME or CE for one pathologist or laboratory professional
- NMB1 Reporting option with CME or CE credit for each additional pathologist or laboratory professional (within the same institution); must order in conjunction with program NMBA
- Two mailings per year with two cases each mailing
- Earn a maximum of four CME credits (AMA PRA Category 1 Credits™) per pathologist and a maximum of four CE credits per laboratory professional per year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



Immunohistochemistry Prognostic Markers

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

c-Myc/Bcl-2 Immunohistochemistry Tissue Microarray MYCB			
Analyte Program Code Challenges per Shipment			
	МҮСВ		
с-Мус		10	
Bcl-2		10	

This program assesses the laboratory's ability to detect c-Myc and Bcl-2-positivity in large B-cell lymphomas, which have emerged as critical prognostic markers.

Program Information

- Two 10-core tissue microarray slides, one for c-Myc and one for Bcl-2
- Two shipments per year

p16 Immunohistochemistry Tissue Microarray P16 Analyte Program Code Challenges per Shipment P16

neck tumors.

Program Information

- One 10-core tissue microarray slide
- Two shipments per year

Ki-67 Immunohistochemistry Tissue Microarray KI67			
Procedure	Program Code Challenges per Shipment		
	KI67		
Ki-67		10	

The purpose of this program is to assess the laboratory's ability to accurately quantify the Ki-67 proliferation index, which is prognostically significant and emerging as a companion diagnostic.

Program Information

- One 10-core cell line tissue microarray slide
- · Two shipments per year

These immunohistochemistry programs assess instrument analytic and pathologist readout steps. For programs focusing on preanalytic steps, see the HistoQIP IHC programs on pages 287-294.

Specialty Anatomic Pathology

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Autopsy Pathology AUP/AUP1			
Procedure	Program Code Challenges per Shipment		
	AUP/AUP1		
Autopsy online case analysis	I	5	

- Program AUP prepares pathologists and pathologists' assistants to succeed by providing ongoing diagnostic learning in autopsy pathology.
- Each case includes case description, gross and/or microscopic images, and case discussion with sample death certificate, key teaching points, and current references.

- AUP Online activity
 providing five cases and the
 second activity includes an
 additional mini-symposium;
 reporting with CME or CE
 credit is available for one
 pathologist or pathologists'
 assistant; for each additional
 pathologist/pathologists'
 assistant, order AUP1
- Includes the option to download program content
- AUP1 Reporting option with CME or CE credit for each additional pathologist or pathologists' assistant (within the same institution); must order in conjunction with program AUP
- Earn a maximum of 12.5 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 12.5 CE credits per pathologists' assistant for completion of entire year.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Online whole slide images powered by DigitalScope technology (if available)
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



Digital Slide Program—Dermatopathology DPATH/DPATH1			
Program Code Challenges per Shipment			
	DPATH/DPATH1		
Online dermatopathology case review 6			

- Program DPATH prepares pathologists, dermatopathologists, and dermatologists to succeed by providing ongoing diagnostic learning in dermatopathology.
- · Cases include static images.
- See system requirements on page 12.

- DPATH Six diagnostic challenges/whole slide images with clinical history; reporting with CME credit is available for one pathologist; for each additional pathologist, order DPATH1
- DPATH1 Reporting option with CME credit for each additional pathologist (within the same institution); must order in conjunction with program DPATH
- · Earn a maximum of 15 CME credits (AMA PRA Category 1 Credits) per pathologist for completion of an entire year.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Hematopathology Online Education HPATH/HPATH1				
Program Code Challenges per Shipment				
HPATH/HPATH1				
Hematopathology online case review	ı	5		

HPATH/HPATH1 prepares pathologists and pathology trainees with an interest in hematopathology to succeed by providing ongoing diagnostic learning and self-assessment.

The program includes 10 challenge cases per year written by expert hematopathologists. For each case the learner will:

- Review virtual whole slide image(s) in the context of a clinical scenario and relevant laboratory data.
- Select and interpret appropriate ancillary studies (IHC, flow cytometry, FISH, cytogenetics, molecular studies, etc) with real-time feedback.
- Simulate a real-world diagnostic workup to arrive at the appropriate diagnosis.
- Apply knowledge to answer three CME questions.

Cases span the breadth of benign and neoplastic hematopathology. Discussions incorporate current best practices in the workup, diagnosis, and subclassification of hematolymphoid diseases.

Neoplastic entities are discussed in the context of both the International Consensus Classification (ICC) and the fifth edition of the World Health Organization (WHO) classification.

See system requirements on page 12.

- HPATH Five diagnostic challenges per activity (two activities per year), with online whole slide images. Reporting with CME credit is available for one participant.
- HPATH1 Reporting option with CME credit for each additional participant (within the same institution); must order in conjunction with program HPATH
- Earn a maximum of 12.5 CME credits (AMA PRA Category 1 Credits™) per participant.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified <u>via email</u> when the activity is available



Neuropathology Program NP/NP1		
Program	Program Code	Challenges per Shipment
	NP/NP1	
Neuropathology online case review	I	8

Program NP prepares anatomic pathologists, neuropathologists, and trainees to succeed by providing ongoing diagnostic learning in neuropathology. Each shipment of this educational program includes eight cases that cover the spectrum of neoplastic and nonneoplastic disorders affecting the central and peripheral nervous systems, including infectious, degenerative, developmental, demyelinating, traumatic, toxicmetabolic, vascular, and neuromuscular diseases. In addition, each mailing will include a mini-symposium focusing on a specific problem area in neuropathology that relates to at least four of the eight cases.

- NP Online activity providing eight cases and a minisymposium; reporting with CME credit is available for one pathologist; for each additional pathologist, order NP1
- Includes option to download program content
- NP1 Reporting option with CME credit for each additional pathologist (within the same institution); must order in conjunction with program NP
- · Earn a maximum of 10 CME credits (AMA PRA Category 1 Credits) per pathologist.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Cytopathology

Analytes/procedures in bold type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Glass Slide Gynecologic Cytopathology PT Program With Glass Slide PAP Education PAP PT

Slide Type		Program Code				Challenges per Year	
	PAPCPT	PAPKPT	PAPMPT	PAPLPT	PAPJPT	Proficiency Testing	Education
Conventional	•				ı	10	10
SurePath							
ThinPrep							
Individual Participant Response Form	APAPCPT	APAPKPT	APAPMPT	APAPLPT	APAPJPT		

Programs PAPCPT, PAPKPT, PAPMPT, PAPLPT, and PAPJPT prepare pathologists and cytotechnologists to succeed by providing ongoing diagnostic learning in gynecologic cytopathology.

Ordering Information

You will receive one shipment for proficiency testing (10 slides) and two additional shipments for your education (five slides each).

Follow these steps to order your PAP Proficiency Testing and PAP Education:

- 1. Choose the following:
 - a. Slide type program code (refer to table above)
 - b. PAP Education series shipment dates (choose one)
 - Series 1
 - O A mailing ships in February
 - o B mailing ships in August
 - Series 2
 - o A mailing ships in May
 - B mailing ships in November
 - c. Add the PAP Education series number after the slide type program code (eg, PAPCPT1, PAPCPT2).
- Order one Individual Participant Response Form code for each participating pathologist/cytotechnologist. Include the PAP Education Series number after the program code (eg, APAPCPT1).
- 3. Select one primary testing session option with two alternative date options using the Gynecologic Cytology Proficiency Testing Order Details Form.
- 4. PPTENR is required by CMS as verification that personnel required to participate in PAP PT under its CLIA number are taking the examination at another laboratory.

Additional Information

- Participants will receive an evaluation via email shortly after submitting results.
- The PAP Education component meets the CAP Laboratory Accreditation Program requirement for participation in a peer educational program.

- Ten glass slides for proficiency testing and 10 glass slides for education
- APAPCPT, APAPKPT, APAPMPT, APAPLPT, APAPJPT - Reporting option with CME or CE credit for each pathologist/ cytotechnologist (within the same institution); must order in conjunction with PAPCPT, PAPKPT, PAPMPT, PAPLPT, PAPJPT
- Earn a maximum of eight CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of eight CE credits per cytotechnologist for completing all challenges.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Three shipments per year; one shipment for proficiency testing (10 slides) and two shipments for education (five slides each)



Cytopathology Glass Slide Education Program PAPCE, PAPJE, PAPKE, PAPLE, PAPME Series 1 or 2

Slide Type		Program Code			Education Challenges per Year	
	PAPCE	PAPKE	PAPME	PAPLE	PAPJE	
Conventional	•				•	
SurePath						
ThinPrep						10
Individual Participant Response Form	APAPCE	APAPKE	APAPME	APAPLE	APAPJE	10

Programs PAPCE, PAPKE, PAPME, PAPLE, and PAPJE prepare pathologists and cytotechnologists to succeed by providing ongoing diagnostic learning in cytopathology.

Ordering Information

Follow these steps to order your PAP Education:

- 1. Choose the following:
 - a. Slide type program code (refer to table above)
 - b. PAP Education series shipment dates (choose one)
 - Series 1
 - O A mailing ships in February
 - O B mailing ships in August
 - Series 2
 - o A mailing ships in May
 - o B mailing ships in November
 - c. Add the PAP Education series number after the slide type program code (eg, PAPCE1, PAPCE2).
- 2. Order one Individual Participant Response Form code for each participating pathologist/cytotechnologist. Include the PAP Education series number after the program code (eg, APAPCE1).

Additional Information

- Participants will receive an evaluation via email shortly after submitting the laboratory form via fax.
- The PAP Education component meets the CAP Laboratory Accreditation Program requirement for participation in a peer educational program.

- · Ten glass slides for education
- APAPCE, APAPJE, APAPKE, APAPLE, APAPME - Reporting option with CME or CE credit for each pathologist/ cytotechnologist (within the same institution); must order in conjunction with programs PAPCE, PAPJE, PAPKE, PAPLE, PAPME
- · Earn a maximum of eight CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of eight CE credits per cytotechnologist for completing all challenges.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Two shipments (five slides each)



Human Papillomavirus (High Risk) for Cytopathology CHPVD, CHPVM, CHPVK, CHPVJ

Analyte/Procedure		Program Code		Challenges per Shipment	
	CHPVD	СНРУМ	CHPVK	CHPVJ	
HPV	ı	•	I	I	5
High-risk HPV genotyping (optional)					5

Additional Information

- Each laboratory should choose the program that best reflects the transport media received in its facility. For program CHPVJ, participants must provide results for all three media types. If your laboratory receives only two types of media, order the programs that are appropriate for your specific laboratory (CHPVD, CHPVM, or CHPVK).
- For laboratories that perform HPV genotyping using ThinPrep PreservCyt or SurePath Preservative Fluid transport mediums on site, programs CHPVM, CHPVK, and select CHPVJ specimens provide an opportunity to report specific HPV genotypes.
- The CAP does not report genotyping responses to the CMS.

Program Information

- Five simulated cervical specimens
- CHPVD Digene® Specimen Transport Medium™ (STM)
- CHPVM ThinPrep PreservCyt® transport medium
- CHPVK SurePath™
 Preservative Fluid transport medium and corresponding vial of diluent
- CHPVJ Combination of Digene, ThinPrep PreservCyt, and SurePath transport mediums
- · Three shipments per year

Color Atlas of Hematology—Vol 1. Peripheral Blood Color Atlas of Hematology—Vol 2. Bone Marrow

The second edition of *Color Atlas of Hematology* has now expanded to two volumes, with the addition of bone marrow pathology.

Volume 1 presents keen insights into peripheral blood pathology with links to 18 engaging videos. View 100+ peripheral blood smears online with DigitalScope® technology.

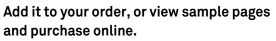
Volume 2 is a useful and instructional reference guide to bone marrow pathology. Explore the detailed "A Closer Look At..." sections. Access the links to interactive slide images.

Vol 1. Peripheral Blood

Item number: PUB222 Hardcover; 480 pages; 2018

Vol 2. Bone Marrow

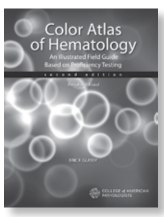
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Touch Imprint/Crush Preparation TICP/TICP1		
Procedure	Program Code	Challenges per Shipment
	TICP/TICP1	
Online slide and image program in rapid assessment case review		4

- The TICP program gets surgical pathologists, cytopathologists, and cytotechnologists ready to succeed by familiarizing them with the cytomorphologic features of pathologic processes and tumors in touch imprints and crush or scrape preparations. These specimens are prepared either for intraoperative consultation (frozen section) or rapid on-site evaluation (ROSE) of tissue biopsies for adequacy and/or interpretation. Participants will learn to make an immediate adequacy assessment, assign the process to a general category, and triage the specimen to appropriate ancillary studies. Participants will review digital whole slides of the TICP preparations (hematoxylin & eosin, modified Wright-Giemsa, and/or Papanicolaou stains), static images of the preparation and ancillary studies, and clinical history/radiographic findings to reach a diagnosis. Each case has a complete description of entities in the differential diagnosis along with a discussion of the correct interpretation.
- Participants will receive immediate feedback on interpretations, ancillary studies, and case-related adequate assessment.
- · The cases will focus on miscellaneous topics.
- May include rarely captured cases that may not be available on the glass slide
- See system requirements on page 12.

- TICP Four online assessment challenges with clinical history; TICP provides CME or CE credit for one pathologist or cytotechnologist; for each additional pathologist or cytotechnologist, order TICP1
- TICP1 Reporting option with CME or CE credit for each additional pathologist/ cytotechnologist (within the same institution); must order in conjunction with program TICP
- · Earn a maximum of 10 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 10 CE credits per cytotechnologist for completion of an entire year.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Online whole slide images powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Nongynecologic Cytopathology Education Program NGC/NGC1			
Procedure	Program Code	Challenges per Shipment	
	NGC/NGC1		
Nongynecologic cytopathology case review – glass slides	ı	5	
Nongynecologic cytopathology case review – online	ı	5 per year	

- Designed to help pathologists and cytotechnologists get ready to succeed, the
 Nongynecologic Cytopathology Education Program (NGC) is an interlaboratory
 educational opportunity to assess participants' screening and interpretive skills.
 The NGC program is unsuitable for proficiency testing as these cases are chosen
 for their educational value. Cases may incorporate static online images that include
 radiology and multiple aspects of pathology to enhance the interpretation.
- Participants will receive an evaluation <u>via email</u> shortly after submitting the laboratory form via fax.
- Additional online advanced education cases provide immediate feedback on interpretation selection, follow-up recommendations, and case-related educational questions.
- See system requirements on page 12.

- NGC Five glass slides per shipment; five online cases; one laboratory response form and two individual response forms
- NGC1 Reporting option with CME or CE credit for each additional pathologist/ cytotechnologist (within the same institution); must order in conjunction with program NGC
- Earn a maximum of 25 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 25 CE credits per cytotechnologist for completing the glass slides and online cases.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- One complimentary online activity with whole slide images powered by DigitalScope technology
- Four shipments of glass slides per year



Digital Slide Program in Fine-Needle Aspiration FNA/FNA1			
Procedure	Program Code	Challenges per Shipment	
	FNA/FNA1		
Online program in fine-needle aspiration case review	ı	5	

- The FNA program gets pathologists and cytotechnologists ready to succeed by focusing on fine-needle aspiration diagnostic dilemmas in practice. Online cases, which consist of whole slide images and static images, provide immediate feedback on interpretation selection, ancillary studies selection, and case-related educational questions.
- Cases will focus on fine-needle aspiration of pancreas and lung topics.
- May include rarely captured cases that may not be available on the glass slide
- See system requirements on page 12.

- FNA Five online diagnostic challenges; FNA provides CME or CE credit for one pathologist or cytotechnologist; for each additional pathologist or cytotechnologist, order FNA1
- FNA1 Reporting option with CME or CE credit for each additional pathologist/ cytotechnologist (within the same institution); must order in conjunction with program FNA
- · Earn a maximum of 10 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 10 CE credits per cytotechnologist.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Online whole slide images powered by DigitalScope technology
- Two online activities per year; your CAP shipping contact will be notified via email when the activity is available

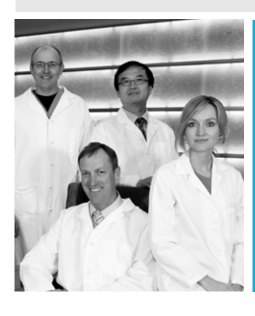


Fine-Needle Aspiration Glass Slide		FNAG/FNAG1	
Procedure	Program Code	Challenges per Shipment	
	FNAG/FNAG1		
Fine-needle aspiration glass slide case review	ı	5	

- The Fine-Needle Aspiration Glass Slide program gets pathologists and cytotechnologists ready to succeed through an interlaboratory educational opportunity to assess participants' screening and interpretive skills. Program FNAG cases may include more than one slide of varying stains and/or preparations used on fine-needle aspirations.
- Cases may include static online images that incorporate radiology and multiple aspects of pathology to support the interpretation.
- Participants will receive an evaluation <u>via email</u> shortly after submitting the laboratory form via fax.

- FNAG Five cases consisting of glass slides and selected online images, representing a variety of conditions; one laboratory response form and two individual response forms
- FNAG1 Reporting option with CME or CE credit for each additional pathologist/ cytotechnologist (within the same institution); must order in conjunction with program FNAG
- Earn a maximum of 10 CME credits (AMA PRA Category 1 Credits) per pathologist and a maximum of 10 CE credits per cytotechnologist.
- This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- Two shipments per year





Benefit from the support of experts in laboratory medicine.

These experts spend countless hours monitoring testing trends to:

- Determine specimen specifications for PT programs to challenge participants.
- Keep our offerings contemporary with new analytes and programs.
- Provide peer-reviewed continuing medical education, continuing education, and self-assessment modules.

Forensic Sciences

Analytes/procedures in **bold** type are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Forensic Pathology FR/FR1			
Procedure	Program Code Challenges per Shipment		
	FR/FR1		
Forensic pathology cases		5	

Additional Information

- Cases may include or reflect anthropologic materials, ballistics, dental identification, DNA identification, environmental pathology, forensic evidence, injury pattern, medicolegal issues, toxicology, and trace evidence.
- FR prepares hospital-based pathologists, forensic pathologists, residents, fellows, and medical examiners/coroners for success by keeping them current in forensic pathology techniques and practices. This educational program is also designed for investigators, analysts, and technicians/technologists.

- FR Online activity containing five case studies illustrating gross and/or microscopic slides and questions related to medicolegal decision making; CME or CE credit is available for one pathologist or investigator. For each additional pathologist or investigator, order FR1.
- FR1 Additional pathologist or investigator (within the same institution) reporting option with CME or CE credit; must order in conjunction with program FR
- · Includes option to download program content
- · Earn a maximum of 12.5 CME credits (AMA PRA Category 1 Credits™) per pathologist and a maximum of 12.5 CE credits per investigator for completion of an entire year.
- · This activity meets the ABPath CC requirements for Improvement in Health and Health Care (IHHC).
- · Two online activities per year; your CAP shipping contact will be notified via email when the activity is available



Vitreous Fluid, Postmortem VF			
Analyte	Program Code	Challenges per Shipment	
	VF		
Acetone		3	
Chloride		3	
Creatinine		3	
Ethanol		3	
Glucose		3	
Potassium		3	
Sodium	I	3	
Vitreous urea nitrogen		3	

Program Information

- Three 5.0-mL synthetic vitreous fluid specimens
- For forensic and other toxicology laboratories that perform quantitative analysis of vitreous fluid
- Conventional and International System of Units (SI) reporting offered
- Two shipments per year



Clinical Toxicology Testing: A Guide for Laboratory Professionals, Second Edition

This book is a practical guide to directing hospital toxicology laboratory operations. This edition features expanded sections on testing in the clinical setting, methodologies, and more user-friendly information on specific analytes. It provides the reader with a comprehensive view of what is needed—and expected—when offering a clinical toxicology service.

Contents include:

- Toxicology testing in the clinical setting, including new chapters on pediatric testing and chronic opioid therapy
- Toxicokinetics and methodologies, with new and expanded information on laboratory-developed tests, screening assays, targeted tests, and oral fluids and alternative matrices
- Specific analytes, including novel psychoactive substances and the use of medical cannabis
- Appendices on such useful topics as urine and serum screens, therapeutic drug monitoring, and proficiency testing

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Item number: PUB227 Softcover; 368 pages; 2020

Forensic Toxicology, Criminalistics FTC			
Analyte	Program Code	Challenges per Shipment	
	FTC		
See drug listing below	I	5	

Program Information

- Five 20.0-mL whole blood specimens
- For crime and hospital laboratories that have forensic toxicology divisions performing qualitative and quantitative analysis of drugs in whole blood specimens
- Three shipments per year

FTC Program Drug Listing

Challenges will include a mix of drugs from the list below.

6-acetylmorphine (6-AM)	Desmethylsertraline	Methylenedioxyamphetamine	Oxymorphone
7-aminoclonazepam	Dextromethorphan	(MDA)	Paroxetine
7-aminoflunitrazepam	Diazepam	Methylenedioxymethamphetamine	Pentobarbital
7-hydroxymitragynine	Dihydrocodeine	(MDMA)	Phencyclidine
Acetaminophen	Diltiazem	Methylenedioxypyrovalerone	Phenethylamine
Alpha-hydroxyalprazolam	Diphenhydramine	(MDPV)	Pheniramine
Alprazolam	Doxepin	Methylphenidate Metaprolel	Phenobarbital
Amitriptyline	Doxylamine	Metoprolol Midazolam	Phentermine
Amphetamine	Duloxetine	Mirtazapine	Phenylephrine
Aripiprazole	Ecgonine ethyl ester	Mitragynine (Kratom)	Phenytoin
Atenolol	Ecgonine methyl ester	Morphine*	Pregabalin
Atropine	Ephedrine	N-desmethyltramadol	Propoxyphene
Benzoylecgonine	Fentanyl*	Naproxen	Propranolol
Brompheniramine	Flunitrazepam	Norbuprenorphine	Pseudoephedrine
Buprenorphine	Fluoxetine	Norchlordiazepoxide	Quetiapine
Bupropion	Gabapentin	Norclomipramine	Quinine
Butalbital	Gamma-hydroxybutyrate (GHB)	Norcodeine	Ranitidine
Carbamazepine	Hydrocodone	Norcyclobenzaprine	Ritalinic acid
Carbamazepine-10,	Hydromorphone	Nordiazepam	Salicylate
11-epoxide	Hydroxybupropion	Nordoxepin	Sertraline
Carisoprodol	Hydroxyzine	Norfentanyl	Strychnine
Chlordiazepoxide	Ibuprofen	Norfluoxetine	Tapentadol
Chlorpheniramine	Imipramine	Norketamine	Temazepam
Citalopram	Ketamine	Normeperidine	Topiramate
Clomipramine	Lamotrigine	Normirtazapine	Tramadol
Clonazepam	Levetiracetam	Noroxycodone	Trazodone
Clozapine	Lidocaine	Norpropoxyphene	Trimipramine
Cocaethylene	Lorazepam	Norsertraline	Valproic acid
Cocaine	Lysergic acid diethylamide (LSD)	Nortrimipramine	Venlafaxine
Codeine	Meperidine*	Nortriptyline	Verapamil
Cyclobenzaprine*	Mephedrone	Norverapamil	Zolpidem
Delta-9-THC	Meprobamate	O-desmethyltramadol	
Delta-9-THC-COOH	Methadone	Olanzapine	
Demoxepam	Methadone metabolite (EDDP)	Oxazepam	
Desipramine Desmothyleleminramine	Methamphetamine	Oxycodone	*and/or metabolite
Desmethylclomipramine			"anu/or metabolite

22 Analyte/Procedure Index



Performance Analytics Dashboard: Bringing it all together

The complimentary dashboard helps you monitor your CAP PT/EQA and accreditation performance.

- Access all graded PT/EQA result forms, evaluations, and participant summaries from one location.
- Benchmark your laboratory against your peers' and CAP-wide performance.
- View performance to quickly identify trends/patterns to mitigate risk.

Analyte/Procedure Index

The following Analyte/Procedure Index is a comprehensive listing of analytes and corresponding CAP program options. It also includes Calibration Verification/Linearity (CVL) and Quality Cross Check (QCC) programs.

Analytes/procedures in bold type whose corresponding program codes are bold are regulated for proficiency testing by the Centers for Medicare & Medicaid Services (CMS).

Laboratories must perform five challenges three times per year (as noted by boldface) for analytes that are regulated by the CMS.

The X in the LAP ENR column denotes the CAP programs that can be used to fulfill the proficiency testing enrollment requirements for CAP-accredited laboratories. Use this index to identify the correct PT programs that match up to your laboratory's activity menu to meet accreditation requirements. For CAP-accredited laboratories outside the US, enrollment in CAP PT/EQA is required for all tests/activities if a program is available. Refer to program descriptions in this catalog to determine compatibility with your specific methodologies.

Analyte/Procedure	LAP ENR	Program Code	Description	Page
1,25-dihydroxy vitamin D		BMV1	Bone Markers and Vitamins	86
1,5-anhydroglucitol		AG	1,5-Anhydroglucitol	71
3-methoxytyramine		N	Urine Chemistry-Special	69
4-hydroxytriazolam		DFC	Drug-Facilitated Crime	111
5-hydroxyindoleacetic acid, qualitative		N	Urine Chemistry-Special	69
5-hydroxyindoleacetic acid, quantitative	Х	N	Urine Chemistry-Special	69
6-acetylmorphine (6-AM)		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
7-aminoclonazepam		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
7-aminoflunitrazepam		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
7-hydroxymitragynine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
11-deoxycortisol		Y/YY	Sex Hormones	84
11-hydroxy-THC		THCB	Blood Cannabinoids	109
17-hydroxycorticosteroids		N	Urine Chemistry-Special	69

Analyte/Procedure	LAP ENR		Description	Page
17-hydroxyprogesterone	X	Y/YY	Sex Hormones	84
17-ketosteroids		N	Urine Chemistry-Special	69
25-OH vitamin D, total	Х	ABVD	Accuracy-Based Vitamin D	114
		LN40	Vitamin D CVL	134
	Х	VITD	25-OH Vitamin D	84
50:50 mixing study, aPTT		CGE/CGEX	Coagulation, Extended	167
		CGS1	Coag Special, Series 1	168
50:50 mixing study, PT		CGE/CGEX	Coagulation, Extended	167
		CGS1	Coag Special, Series 1	168
ABO grouping	Х	J, JXM, J1	Transfusion Medicine	232
	Х	JAT, JATXM	Transfusion Medicine, Automated	233
		JATE1	Transfusion Medicine, Automated, Educational	233
		JATQ	QCC, Transfusion Medicine	48
		TMCA	Transfusion Medicine, Competency Assessment	239
ABO subgroup typing		ABOSG	ABO Subgroup Typing	235
		J, JXM	Transfusion Medicine	232
		JAT, JATXM	Transfusion Medicine, Automated	233
Acetaminophen	Х	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FTC	Forensic Toxicology, Criminalistics	107
		LN3	TDM CVL	125
		SDS	Serum Drug Screen	104
		T	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Acetone	Х	AL1	Whole Blood Alcohol/ Volatiles	104
	Х	AL2	Serum Alcohol/Volatiles	104
		SDS	Serum Drug Screen	104

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Acetone (cont.)		VF	Vitreous Fluid, Postmortem	104
Acid phosphatase		C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Acid-fast smear	X	E	Mycobacteriology	193
	X	E1	Mycobacteriology, Ltd	193
Acinetobacter calcoaceticus- baumannii complex	X	IDPN	Infectious Disease, Pneumonia Panel	211
Activated clotting time	Х	CT, CT1, CT2, CT3, CT5	ACT	170
		CTQ, CT1Q, CT2Q, CT3Q, CT5Q	QCC, ACT	46
		POC14, POC15, POC16	Competency Activated Clotting Time	52
Activated partial thromboplastin time		APXBN	Anticoagulant Monitoring, Apixaban	170
	Х	CGB	Basic Coagulation	166
		CGE/CGEX	Coagulation, Extended	167
	X	CGL	Coagulation, Limited	166
		CGLQ	QCC, Coagulation, Limited	46
		CGS1	Coag Special, Series 1	168
		CGS3	Coag Special, Series 3	168
		CGS4	Coag Special, Series 4	168
		DBGN	Anticoagulant Monitoring, Dabigatran	170
		FNPX	Anticoagulant Monitoring, Fondaparinux	170
		RVBN	Anticoagulant Monitoring, Rivaroxaban	170
Activated protein C resistance		CGE/CGEX	Coagulation, Extended	167
		CGS2	Coag Special, Series 2	168
Active vitamin B ₁₂		MMA	MMA and Active Vitamin	82
Acylcarnitine		BGL	Biochemical Genetics	257
Acylcarnitine quantitation		BGL4	Acylcarnitine Quantitation for Inherited Metabolic Disorders	259
ADAMTS13		CGS7	ADAMTS13	168
Adenovirus		GIP	Gastrointestinal Panel	212
	X	GIP5	Gastrointestinal Panel, 5 Challenge	212
		GIPN	Gastrointestinal Panel, Global	213

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Adenovirus (cont.)		ID2	Nucleic Acid Amp,	204
			Respiratory	
	Х	IDPN	Infectious Disease, Pneumonia Panel	211
	Х	IDR	Infectious Disease, Respiratory Panel	210
		VLS2	Viral Load	206
	Х	VR1	Virology Culture	200
	Х	VR2	Viral Antigen by DFA	200
	X	VR4	Viral Antigen by EIA and Latex	200
Adenovirus 40/41		SP, SPN	Stool Pathogen	189
Adrenocorticotropic hormone (ACTH)	X	TM/TMX	Tumor Markers	89
Alanine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Alanine aminotransferase (ALT/SGPT)	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Albumin		ABS	Accuracy-Based Testosterone and Estradiol	115
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
		SPE	Protein Electrophoresis	76
Albumin, CSF	Х	M, OLI	CSF Chemistry and Oligoclonal Bands	74
Albumin, urine		ABU	Accuracy-Based Urine	115
		LN20	Urine Albumin	129
	Х	U	Urine Chemistry-General	68
Albumin:creatinine ratio		ABU	Accuracy-Based Urine	115

Analyte/Procedure	LAP ENR	0	Description	Page
Albumin:creatinine ratio (cont.)		LN20	Urine Albumin CVL	129
		U	Urine Chemistry-General	68
		UMC	Urine Albumin Creatinine	160
Alcohol, serum	Х	AL2	Serum Alcohol/Volatiles	104
		LN11	Serum Ethanol CVL	127
Alcohol, whole blood	X	AL1	Whole Blood Alcohol/ Volatiles	104
Aldolase		ADL	Aldolase	71
Aldosterone, serum	X	RAP	Renin and Aldosterone	89
Aldosterone, urine		N	Urine Chemistry-Special	69
Alkaline phosphatase (ALP)	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD2	Body Fluid Chemistry 2	73
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Allergens (specific)		SE	Diagnostic Allergy	221
Alloisoleucine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Alpha-1 antitrypsin	Х	IG/IGX	Immunology, General	216
		LN7	Immunology CVL	126
Alpha-1 antitrypsin genotyping (SERPINA1) gene	X	AAT	Alpha-1 Antitrypsin Genotyping	259
Alpha-1 globulin		SPE	Protein Electrophoresis	76
Alpha-2 globulin		SPE	Protein Electrophoresis	76
Alpha-2-antiplasmin		CGE/CGEX	Coagulation, Extended	167
Alpha-2-macroglobulin		A2MG	Alpha-2-Macroglobulin	218
Alpha-fetoprotein (AFP), amniotic fluid	Х	FP/FPX	Maternal Screen	87
Alpha-fetoprotein (AFP), serum	Х	FP/FPX	Maternal Screen	87
	Χ	K/KK	Ligand-General	82
		LN5	Ligand CVL	125
		LN5S	Ligand CVL – all Siemens ADVIA (Centaur, CP, and XP) and Atellica IM	125
Alpha-hydroxyalprazolam		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107

Analyte/Procedure	LAP ENR		Description	Page
Alpha-hydroxyalprazolam (cont.)		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Alpha-thalassemia		HGM	Hemoglobinopathies, Molecular Methods	261
Alprazolam		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UT	Urine Toxicology	98
Aluminum	X	R	Trace Metals	78
Aluminum, urine		TMU	Trace Metals, Urine	106
Aluminum, whole blood		TMWB	Trace Metals, Whole Blood	106
Amikacin	Х	CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN3	TDM CVL	125
Amino acids, qualitative	X	BGL	Biochemical Genetics	257
Amino acids, quantitative		BGL	Biochemical Genetics	257
		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Amitriptyline		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
	Х	ZT	TDM, Special	59
Ammonia		C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN32	Ammonia CVL	132
Amniotic fluid leakage (nitrazine)		AFL	Amniotic Fluid Leakage	154
Amobarbital		DFC	Drug-Facilitated Crime	111
Amphetamine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98

Analyte/Procedure		Program Code	Description	Page
	EINK	Code		
Amphetamine (cont.)		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
		UTCO	Urine Toxicology Carryover	137
Amphetamine group		DMPM	Drug Monitoring for Pain Management	110
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Amylase	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Amylase, pancreatic	Х	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Amylase, urine		LN6	Urine Chemistry CVL	126
	Х	U	Urine Chemistry-General	68
Anaerococcus prevotii/ vaginalis		JIP	Joint Infection Panel	208
Anaplasma phagocytophilum		TTD	Antibody Detection of Tick-Transmitted Diseases	214
Anaplastic lymphoma kinase	Х	PM6	Anaplastic Lymphoma Kinase IHC	298
Androstenedione	Х	Y/YY	Sex Hormones	84
Angiotensin converting enzyme		ACE	Angiotensin Converting Enzyme	71
Anti ADAMTS13 IgG		CGS7	ADAMTS13	168
Anti-A titer		AABT, AABT1	Antibody Titer, Automated	237
		ABT, ABT1	Antibody Titer	236
Anti-B titer		AABT3	Antibody Titer, Automated	237
		ABT3	Antibody Titer	236
Antibody detection	Х	J, JXM, JAT,	Transfusion Medicine	232-
		JATXM		233

Analyte/Procedure	LAP ENR	0	Description	Page
Antibody detection (cont.)		JATE1	Transfusion Medicine, Automated, Educational	233
		JATQ	QCC, Transfusion Medicine	48
	Χ	PS	Platelet Serology	238
		TMCA	Transfusion Medicine, Competency Assessment	239
Antibody detection/ identification (HLA)	Х	MXC, MXEP	HLA Analysis, Class I/II	248
Antibody identification		ETME1	Expanded Transfusion Medicine Exercises	242
	Х	J, JXM, JAT, JATXM	Transfusion Medicine	232- 233
		JATE1	Transfusion Medicine, Automated, Educational	233
		TMCA	Transfusion Medicine, Competency Assessment	239
Antibody screen (HLA)		MXC, MXEP, MXS	HLA Analysis, Class I/II	248
Antibody titer		ABT, ABT1, ABT2, ABT3	Antibody Titer	236
Antibody titer, automated		AABT, AABT1, AABT2, AABT3	Antibody Titer, Automated	237
Anticardiolipin IgA, qualitative		ACL, APS	Antiphospholipid Antibody	219
Anticardiolipin IgA, quantitative		ACL, APS	Antiphospholipid Antibody	219
Anticardiolipin IgG, IgM, polyclonal; qualitative	Х	ACL, APS	Antiphospholipid Antibody	219
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		UT	Urine Toxicology	98
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		OFD	Oral Fluid for Drugs of Abuse	103
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Doxepin		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98

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		UT	Urine Toxicology	98
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		UT	Urine Toxicology	98
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Ecgonine methyl ester		FTC	Forensic Toxicology, Criminalistics	107
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		UT	Urine Toxicology	98
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		GIPN	Gastrointestinal Panel, Global	213
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	ENR	Code		
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		GIPN	Gastrointestinal Panel, Global	213
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		IDME	Meningitis/Encephalitis Panel	209
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	X	IDM5	Meningitis/Encephalitis Panel	209
Escherichia coli 0157		GIP	Gastrointestinal Panel	212
	Х	GIP5	Gastrointestinal Panel, 5 Challenge	212
Estazolam		DFC	Drug-Facilitated Crime	111
Estradiol		ABS	Accuracy-Based Testosterone and Estradiol	115
		LN8	Reproductive Endocrinology CVL	127
	Х	Y/YY	Sex Hormones	84
Estriol, unconjugated (uE3)	X	FP/FPX	Maternal Screen	87
	X	Y/YY	Sex Hormones	84
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Ethanol	X	AL1	Whole Blood Alcohol/ Volatiles	104
	X	AL2	Serum Alcohol/Volatiles	104
		LN11	Serum Ethanol CVL	127
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Ethanol, vitreous fluid		VF	Vitreous Fluid, Postmortem	104
Ethosuximide		CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Ethyl glucuronide (EtG)		ETB	Ethanol Biomarkers	105
Ethyl sulfate (EtS)		ETB	Ethanol Biomarkers	105
Ethylene glycol		AL1	Whole Blood Alcohol/ Volatiles	104
		AL2	Serum Alcohol/Volatiles	104
Etizolam		DFC	Drug-Facilitated Crime	111
		T	Toxicology	98
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	X	TPM	Thrombophilia Mutations	265
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		ECF	Expanded Coagulation Factors	167

Analyte/Procedure	LAP ENR	Program Code	Description	Page
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	Х	TPM	Thrombophilia Mutations	265
Factor VII		CGE/CGEX	Coagulation, Extended	167
		ECF	Expanded Coagulation Factors	167
Factor VIII		CGE/CGEX	Coagulation, Extended	167
		CGS3	Coag Special, Series 3	168
		ECF	Expanded Coagulation Factors	167
		LN51	Factor VIII CVL	133
Factor VIII inhibitor		CGS3	Coag Special, Series 3	168
Factor IX		CGE/CGEX	Coagulation, Extended	167
		ECF	Expanded Coagulation Factors	167
Factor X		CGE/CGEX	Coagulation, Extended	167
		ECF	Expanded Coagulation Factors	167
Factor XI		CGE/CGEX	Coagulation, Extended	167
		ECF	Expanded Coagulation Factors	167
Factor XII		CGE/CGEX	Coagulation, Extended	167
		ECF	Expanded Coagulation Factors	167
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		ECF	Expanded Coagulation Factors	167
Familial dysautonomia (ELP1 gene)	X	MGL4	Molecular Genetics	262- 263
Fanconi anemia, complementation grp. C (FANCC gene)	X	MGL4	Molecular Genetics	262- 263
Fecal calprotectin		FCAL	Fecal Calprotectin	75
Fecal fat, qualitative		FCFS	Fecal Fat	75
Fecal lactoferrin		FLAC	Fecal Lactoferrin	187
Fecal occult blood		OCB	Occult Blood	159
		OCBQ	QCC, Occult Blood	45
Fentanyl		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
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		CZQ	QCC, Chemistry and TDM	37
	Х	K/KK	Ligand-General	82
		LN5	Ligand CVL	125
		LN5S	Ligand CVL – all Siemens ADVIA (Centaur, CP, and XP) and Atellica IM	125
Fetal fibronectin	Х	FF	Fetal Fibronectin	88
Fetal hemoglobin (gastric fluid)		APT	Fetal Hemoglobin	157
Fetal hemoglobin identification	Х	HG	Hemoglobinopathy	147
Fetal membrane rupture		ROM1	Fetal Membranes/ Preterm Labor	159
Fetal red cell quantitation	Х	HBF	Fetal Red Cell Detection	238
		TMCAF	Transfusion Medicine, Competency Assessment	239
Fetal screen (Rosette testing)	Х	HBF	Fetal Red Cell Detection	238
		TMCAF	Transfusion Medicine, Competency Assessment	239
Fibrin degradation products, plasma		CGDF	Coagulation, D-dimer/ FDP	166
		CGL	Coagulation, Limited	166
		CGLQ	QCC, Coagulation, Limited	46
Fibrin degradation products, serum		CGDF	Coagulation, D-dimer/ FDP	166
		CGL	Coagulation, Limited	166
		CGLQ	QCC, Coagulation, Limited	46
Fibrin monomer		CGL	Coagulation, Limited	166
		CGDF	Coagulation, D-dimer/ FDP	166
Fibrinogen	X	CGL	Coagulation, Limited	166
		CGLQ	QCC, Coagulation, Limited	46
		LN44	Fibrinogen, CVL	134
Fibrinogen antigen		CGE/CGEX	Coagulation, Extended	167
Finegoldia magna		JIP	Joint Infection Panel	208
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Fine-needle aspiration, glass slides		FNAG/ FNAG1	Fine-Needle Aspiration	312

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FISH for breast carcinoma hybridization and interpretation on site ERBB2 (HER2) amplification	X	СҮН	FISH for ERBB2 (HER2) Amplification	255
FISH for breast carcinoma, interpretation only, <i>ERBB2</i> (<i>HER2</i>) gene amplification		СҮНІ	FISH for ERBB2 (HER2) Amplification, Interpretation Only Exercise	255
FISH for constitutional and hematologic disorders		CYF	Fluorescence In Situ Hybridization and Interpretation on Site	254
FISH for lung cancer, ALK rearrangement		CYALK	Fluorescence In Situ Hybridization and Interpretation on Site, Lung Cancer	255
FISH for lymphoma		CYL	Fluorescence In Situ Hybridization and Interpretation on Site, Lymphoma	255
FISH for paraffin- embedded tissue	Х	CYH	FISH for ERBB2 (HER2) Amplification	255
		CYJ	Fluorescence In Situ Hybridization and Interpretation on Site, Brain/Glioma Tissue	255
		СҮК	Fluorescence In Situ Hybridization and Interpretation on Site, Solid Tumor	255
		CYL	Fluorescence In Situ Hybridization and Interpretation on Site, Lymphoma	255
FISH for solid tumor		СҮК	Fluorescence In Situ Hybridization and Interpretation on Site, Solid Tumor	255
FISH for urothelial carcinoma hybridization and interpretation	X	CYI	Fluorescence In Situ Hybridization and Interpretation on Site, Urothelial Carcinoma	254
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Flow cytometry, post- immunotherapy analysis		FL6	Flow Cytometry, Post-immunotherapy Analysis	225
Fluconazole		AFD	Antifungal Drugs Monitoring	109
Flunitrazepam		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98

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Flunitrazepam (cont.)		UT	Urine Toxicology	98
Fluorescent microscope check		I	Instrumentation	137
Fluoxetine		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Folate, RBC	Х	FOL	RBC Folate	88
Folate, serum	X	K/KK	Ligand-General	82
		LN5	Ligand CVL	125
		LN5S	Ligand CVL – all Siemens ADVIA (Centaur, CP, and XP) and Atellica IM	125
Follicle-stimulating hormone (FSH)		ABS	Accuracy-Based Testosterone, Estradiol	115
		LN8	Reproductive Endocrinology CVL	127
	X	Y/YY	Sex Hormones	84
Fondaparinux		FNPX	Anticoagulant Monitoring, Fondaparinux	170
Forensic pathology		FR/FR1	Forensic Pathology	314
Forensic toxicology		FTC	Forensic Toxicology, Criminalistics	107
Fragile X (<i>FMR1</i> gene)	Х	MGL1	Molecular Genetics	262- 263
Free beta hCG		FP1B	First Trimester Maternal Screening, Free Beta	88
Free Kappa/Lambda ratio		SFLC	Serum Free Light Chains	223
Free testosterone		Υ	Sex Hormones	84
Friedreich ataxia (<i>FXN</i> gene)	Х	MGL2	Molecular Genetics	262- 263
Fructosamine		FT	Fructosamine	75
Fungal culture		CBT	Cord Blood Testing	240
		SCP	Stem Cell Processing	240
Fungal serology		FSER	Fungal Serology	196
Fungus identification	Х	F	Mycology and Aerobic Actinomycetes	194
	Х	F1	Yeast	194
	Х	F3	Candida Culture	195
	Х	IDM5	Infectious Disease Meningitis/Encephalitis	209
	X	MVP	Molecular Vaginal Panel	191
	Х	VS	Vaginitis Screen	190
	Х	YBC	Yeast Blood Culture, Molecular	195
G6PD		PGX1	Pharmacogenetics	264
Gabapentin		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110

Analyte/Procedure	LAP ENR	0	Description	Page
Gabapentin (cont.)		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
		ZE	Therapeutic Drug Monitoring, Extended	59
Galactomannan		FGAL	Galactomannan	195
Gamma globulin		M, OL1	CSF Chemistry	74
		SPE	Serum Electrophoresis	76
Gamma glutamyl transferase (GGT)	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Gamma hydroxybutyrate (GHB)		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
Gardnerella vaginalis, DNA probe	Х	VS	Vaginitis Screen	190
Gastric occult blood		GOCB	Gastric Occult Blood	157
Gastric pH		GOCB	Gastric Occult Blood	157
Gastrin		ABGIC	Accuracy Based Glucose, Insulin and C-peptide	118
	Х	ING	Insulin, Gastrin, C-peptide	86
Gaucher disease (GBA gene)	Х	MGL4	Molecular Genetics	262- 263
GDH antigen		CDF2	Clostridioides (Clostridium) difficile Detection	187
	X	CDF5	Clostridioides (Clostridium) difficile Detection	187
	Х	D	Bacteriology	177
Genomic copy number array		CYCGH	Constitutional Microarray Analysis	256
Gentamicin	Х	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN3	TDM CVL	125
Giardia		GIP	Gastrointestinal Panel	212
	Х	GIP5	Gastrointestinal Panel, 5 Challenge	212

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Giardia immunoassay, preserved specimen	Х	P, P3, P4, P5	Parasitology	197
Giemsa stain	Х	ВР	Blood Parasite	198
	Х	Р	Parasitology	197
Glioma by FISH		CYJ	Fluorescence In Situ Hybridization and Interpretation on Site, Brain/Glioma Tissue	255
Glucose		ABGIC	Accuracy-Based Glucose, Insulin, and C-peptide	118
	Х	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
	X	C1, C3/C3X, C4, CZ/ CZX/CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN13C	Blood Gas CVL	128
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Glucose, CSF	Х	M, OLI	CSF Chemistry and Oligoclonal Bands	74
Glucose, urine	Х	CMP, CMP1	Clinical Microscopy	152
		CMQ	QCC, Urinalysis	44
	Χ	HCC2	Waived Combination	66
		LN6	Urine Chemistry CVL	126
		POC3	POC Urine Dipstick Competency	50
	Χ	U	Urine Chemistry-General	68
Glucose, vitreous fluid		VF	Vitreous Fluid, Postmortem	104
Glucose, whole blood	Х	HCC	Waived Combination	65
		HCC2	Waived Combination	66
	X	LCW	Chemistry-Ltd, Waived	64
		LN17	Whole Blood Glucose CVL	129
		POC2	POC Glucose Competency	50
		POC7	POC/Waived Glucose and Hemoglobin Competency	50

Analyte/Procedure		Program Code	Description	Page
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HSV (see Herpes simplex virus)				
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	Х	HCC2	Waived Combination	66
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		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
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International normalized ratio (INR)	Х	CGB	Basic Coagulation	166
	Х	CGL	Coagulation, Limited	166
		CGS1	Coag Special, Series 1	168
		CGS4	Coag Special, Series 4	168
		POC6	POC PT/INR, CoaguChek XS Plus	50
		WP10	Whole Blood Coagulation	173
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		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		POC10, POC11	POC Competency Blood Gases	51
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		CMQ	QCC, Urinalysis	44
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			Immunohistochemistry TMA	
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		JIP	Joint Infection Panel	208
Klebsiella oxytoca	Х	IDPN	Infectious Disease, Pneumonia Panel	211
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		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
	Х	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		LN13C	Blood Gas CVL	128
		POC10, POC11	POC Competency Blood Gases	51

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Lactate dehydrogenase (LD)	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
		SC0	Serum Carryover	137
Lactate dehydrogenase (LD), CSF	Х	M, OLI	CSF Chemistry and Oligclonal Bands	74
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		CZQ	QCC, Chemistry and TDM	37
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Levetiracetam		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
		ZE	Therapeutic Drug Monitoring, Extended	59
Levorphanol		T	Toxicology	98
		UT	Urine Toxicology	98
Lidocaine	Х	CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FTC	Forensic Toxicology, Criminalistics	107
		LN3	TDM CVL	125
		T	Toxicology	98
		UT	Urine Toxicology	98
Lipase	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD2	Body Fluid Chemistry 2	73
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Lipids		ABL	Accuracy-Based Lipid	114
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56

Analyte/Procedure	LAP ENR		Description	Page
Lipids (cont.)		CZQ	QCC, Chemistry and TDM	37
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Lipoprotein (a)	Х	ABL	Accuracy-Based Lipid	114
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Lipoprotein-associated phospholipase		PLA	Lipoprotein-Associated Phospholipase A ₂	75
Lipoprotein electrophoresis		LPE	Lipoprotein Electrophoresis	76
Listeria monocytogenes		IDME	Meningitis/Encephalitis Panel	209
	Х	IDM5	Meningitis/Encephalitis Panel	209
Lithium	X	C1, C3/C3X, CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN3	TDM CVL	125
Liver-kidney microsomal antibody		LKM	Liver-Kidney Microsomal Antibody	221
Lorazepam		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
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Luteinizing hormone (LH)		ABS	Accuracy-Based Testosterone, Estradiol	115
		LN8	Reproductive Endocrinology CVL	127
	Χ	Y/YY	Sex Hormones	84
Lysine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
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Lysergic acid diethylamide (LSD)		FTC	Forensic Toxicology, Criminalistics	107
		UDS, UDS6	Urine Drug Screen	100
Magnesium	X	C1,C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Magnesium, ionized	X	AQ, AQH	Critical Care Blood Gas	92-93
		AQQ, AQHQ	QCC, Critical Care Blood Gas Series	42
		POC10, POC11	POC Competency Blood Gases	51
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Malaria	Х	RML5	Rapid Malaria, 5 Challenge	198
		RMAL	Rapid Malaria	198
Manganese		R	Trace Metals	78
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Mature B-cell leukemia/ lymphoma measurable (minimal) residual disease		FL8	Flow Cytometry Mature B-cell Leukemia/ Lymphoma Measurable (Minimal) Residual Disease	227
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		HE	Basic Hematology	140
МСНС		FH1-FH4, FH9-FH10, FH13, FH16-FH17	Hematology Automated Differential	140

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MCHC (cont.)		FH3Q, FH4Q, FH9Q, FH13Q	QCC, Automated Hematology Series	43
		HE	Basic Hematology	140
MCV		FH1-FH4, FH9-FH10, FH13, FH16-FH17	Hematology Automated Differential	140
		FH3Q, FH4Q, FH9Q, FH13Q	QCC, Automated Hematology Series	43
		HE	Basic Hematology	140
Measurable (minimal) residual disease		BALL	B-ALL Measurable (Minimal) Residual Disease	227
		FL8	Flow Cytometry Mature B-cell Leukemia/ Lymphoma Measurable (Minimal) Residual Disease	227
		FL9	Flow Cytometry Plasma Cell Myeloma Measurable (Minimal) Residual Disease	228
		MRD	Measurable (Minimal) Residual Disease, <i>BCR/</i> <i>ABL1</i> p210	279
		MRD1	Measurable (Minimal) Residual Disease, <i>BCR/</i> <i>ABL1</i> p190	279
		MRD2	Measurable (Minimal) Residual Disease, <i>PML/</i> <i>RARA</i>	279
MECP2 deletion/ duplication analysis	Х	RETT	Rett Syndrome Genotyping	265
MECP2 genotyping	Х	RETT	Rett Syndrome Genotyping	265
MEN2 (RET gene)	Х	MGL3	Molecular Genetics	262- 263
Meperidine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Mephedrone		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
Manuahamat -		UT	Urine Toxicology	98
Meprobamate		DFC	Drug-Facilitated Crime	111

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Meprobamate (cont.)		DMPM	Drug Monitoring for Pain	110
Meprobamate (cont.)		DIVIFIVI	Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Meprobamate/ Carisoprodol		UDS, UDS6	Urine Drug Screen	100
Mercury, urine		TMU	Trace Metals, Urine	106
Mercury, whole blood		TMWB	Trace Metals, Whole Blood	106
Metabolic disease testing		BGL	Biochemical Genetics	257
Meta- chlorophenylpiperazine (m-CPP)		DFC	Drug-Facilitated Crime	111
		T	Toxicology	98
		UT	Urine Toxicology	98
Metanephrine	X	N	Urine Chemistry-Special	69
Methadone		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Methadone metabolite (EDDP)		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Methamphetamine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98

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Methanol	Х	AL1	Whole Blood Alcohol/ Volatiles	104
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Methaqualone		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
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Methionine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Methicillin-resistant Staphylococcus aureus (MRSA)		BCS1	Blood Culture Staphylococcus aureus	184
		IDN, IDO	Nucleic Acid Amp, Organisms	207
		MRS	Methicillin-Resistant S. aureus Screen	188
		MRS2M	MRSA Screen, Molecular, 2 Challenge	188
	Х	MRS5	Methicillin-Resistant S. aureus Screen	188
	Х	MRS5M	MRSA Screen, Molecular, 5 Challenge	188
Methotrexate	X	CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Methylenedioxy- amphetamine (MDA)		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Methylenedioxyethyl- amphetamine (MDEA)		UDC	Forensic Urine Drug Testing, Confirmatory	102
Methylenedioxymeth- amphetamine (MDMA)		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98

Analyte/Procedure		Program Code	Description	Page
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		T	Toxicology	98
		UT	Urine Toxicology	98
Methylenetetra- hydrofolate reductase (MTHFR gene)		MGL1	Molecular Genetics	262- 263
Methylmalonic acid		MMA	MMA and Active B ₁₂	82
Methylphenidate		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Metoprolol		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
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	Х	U	Urine Chemistry-General	68
	X	UMC	Urine Albumin (Microalbumin)/ Creatinine	160
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Midazolam		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
Mirtazapine		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
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Mitragynine (Kratom)		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
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Mixing studies, PT		CGE/CGEX	Coagulation, Extended	167
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Modified acid-fast stain	Х	P, P3, P4, P5	Parasitology	197
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Molecular genetics	X	MGL1, MGL2, MGL3, MGL4, MGL5	Molecular Genetics	262– 263
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Molecular typing		IDN, IDO	Nucleic Acid Amp, Organisms	207
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Mononuclear cell count		CBT	Cord Blood Testing	240
		SCP	Stem Cell Processing	240
Moraxella catarrhalis	X	IDPN	Infectious Disease, Pneumonia Panel	211
Morganella morganii		JIP	Joint Infection Panel	208
Morphine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
M-protein (paraprotein) identification	X	SPE	Protein Electrophoresis	76
MPL		MH02, MH03	Molecular Hematologic Oncology	278
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Mucolipidosis IV (MCOLN1 gene)	Х	MGL4	Molecular Genetics	262- 263
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Multimodality biomarker assessment		NMBA, NMB1	Navigating Multimodality Biomarker Assessment	300
Multiple endocrine neoplasia type 2 (<i>RET</i> gene)	Х	MGL3	Molecular Genetics	262- 263
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Mycobacterial culture	Х	E1	Mycobacteriology, Ltd	193
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Mycobacterium tuberculosis		IDO	Nucleic Acid Amp, Organisms	207
Mycobacterium tuberculosis antibody detection		QF	M. tuberculosis Infection Detection	221
Mycobacterium tuberculosis identification and resistance detection		MTBR	Molecular MTB Detection and Resistance	193
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Mycoplasma genitalium		MGEN	Mycoplasma genitalium, Molecular	190
		STIM	Sexually Transmitted Infection Detection, Molecular	191
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	Х	IDR	Infectious Disease, Respiratory Panel	210
		VR3	Antibody Detection– Infectious Disease Serology	214
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		CRTQ	QCC, Cardiac Markers	38
		HCRQ	QCC, High-Sensitivity Cardiac Markers	39
		LN33	Serum Myoglobin CVL	132
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		POC12	POC Cardiac Markers Competency	51

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Myotonic dystrophy (DMPK gene)	Х	MGL2	Molecular Genetics	262- 263
N-acetylprocainamide (NAPA)		CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
N-desmethyltramadol		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Naloxone		DMPM	Drug Monitoring for Pain Management	110
Naproxen		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
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Neisseria gonorrhoeae	Х	D3	GC Cultures	179
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	Х	HC7	Nucleic Acid Amp C. trachomatis/GC DNA	191
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		JIP	Joint Infection Panel	208
	Х	RMC	Routine Microbiology Combination	180
	X	STIM	Sexually Transmitted Infection Detection, Molecular	191
Neisseria meningitidis		IDME	Meningitis/Encephalitis Panel	209
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		NGSB1	NGS Solid Tumor Bioinformatics	267
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		NGSB4	NGS Solid Tumor Bioinformatics Hybrid	268
		NGSB5	NGS Hematologic Malignancies Bioinformatics Hybrid	270

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		ТМВ	Tumor Mutational Burden	273
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Nitrite, urine	Х	CMP, CMP1	Clinical Microscopy	152
		CMQ	QCC, Urinalysis	44
		DAI	Urine Drug Adulterant/ Integrity Testing	101
	X	HCC2	Waived Combination	66
		POC3	POC Urine Dipstick Competency	50
Nitrogen, urine; total		U	Urine Chemistry-General	68
Nongynecologic cytopathology		FNA/FNA1	Fine-Needle Aspiration, Digital	311
		FNAG/ FNAG1	Fine-Needle Aspiration, Glass	312
		NGC/NGC1	Nongynecologic Cytopathology Education Program	310
Non-HDL cholesterol, calculated		ABL	Accuracy-Based Lipid	114
Noninvasive prenatal testing		NIPT	Noninvasive Prenatal Testing	88
Norbuprenorphine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Norchlordiazepoxide		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Norclomipramine		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98

Analyte/Procedure	LAP ENR		Description	Page
Norcodeine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Norcyclobenzaprine		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Nordiazepam		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Nordoxepin		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Norepinephrine	X	N	Urine Chemistry-Special	69
Norfentanyl		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Norfluoxetine		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Norhydrocodone		DMPM	Drug Monitoring for Pain Management	110
Norketamine		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Normeperidine		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107

Analyte/Procedure	LAP ENR	0	Description	Page
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		UT	Urine Toxicology	98
Normetanephrine	Х	N	Urine Chemistry-Special	69
Normirtazapine		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Nornaloxone		T	Toxicology	98
		UT	Urine Toxicology	98
Norovirus		GIP	Gastrointestinal Panel	212
	Х	GIP5	Gastrointestinal Panel, 5 Challenge	212
		GIPN	Gastrointestinal Panel, Global	213
		SP1	Stool Pathogens	189
Noroxycodone		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Noroxymorphone		DMPM	Drug Monitoring for Pain Management	110
Norpropoxyphene		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Norsertraline		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Nortrimipramine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Nortriptyline		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
	Х	ZT	TDM, Special	59
Norvenlafaxine		DFC	Drug-Facilitated Crime	111
Norverapamil		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98

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NRAS	Х	MTP	Multigene Tumor Panel	277
nRBC		FH3, FH9, FH13, FH16-FH17	Hematology Automated Differential	140
		FH3Q, FH9Q, FH13Q	QCC, Automated Hematology Series	43
NT-pro B-natriuretic peptides	Х	BNP5	B-type Natriuretic Peptides, 5 Challenge	59
		BNPQ	QCC, B-type Natriuretic Peptides	37
		LN30	BNP CVL	131
	Х	PCARM/ PCARMX	Point-of-Care Cardiac Markers	64
Nucleated cells, total		ABF3	Automated Body Fluid	154
		CBT	Cord Blood Testing	240
		SCP	Stem Cell Processing	240
Nucleated red blood cell count		FH3, FH9, FH13, FH16-FH17	Hematology Automated Differential	140
		FH3Q, FH9Q, FH13Q	QCC, Automated Hematology Series	43
Nucleated red cells, total		CBT	Cord Blood Testing	240
Nucleic acid amplification	Х	HBVL, HBVL5, HCV2	Hepatitis Viral Load	205
	Х	HC6/HC6X	C. trachomatis/GC by Nucleic Acid Amp	191
	Х	НС7	C. trachomatis/GC DNA by NAA	191
	Х	HIVG, HV2	HIV Viral Load	206
		ID1, ID1T	Nucleic Acid Amp, Viruses	201
		ID2	Nucleic Acid Amp, Respiratory	204
	Х	ID3	Nucleic Acid Amplification, Respiratory Limited	204
		ID3Q	QCC, Nucleic Acid Amplification, Respiratory Limited	47
		IDN, IDO	Nucleic Acid Amp, Organisms	207
		MRS2M	MRSA Screen, Molecular, 2 Challenge	188
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		SP, SPN, SP1	Stool Pathogens	189
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Nucleic acid testing	Х	NAT	Nucleic Acid Testing	245
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Nugent scoring		VS2	Vaginitis Screen, Virtual Gram Stain	192
Occult blood		OCB	Occult Blood	159
		OCBQ	QCC, Occult Blood	45
		POC9	POC Fecal Occult Blood	50
Occult blood, gastric		GOCB	Gastric Occult Blood	157
O-desmethyltramadol		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Olanzapine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Oligoclonal bands		OLI	Oligoclonal Bands	74
Opiate group		DMPM	Drug Monitoring for Pain Management	110
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
		UTCO	Urine Toxicology Carryover	137
OPRM1		PGX1	Pharmacogenetics	264
Organic acids, urine; qualitative	Х	BGL	Biochemical Genetics	257
Organic acids, urine; quantitative		BGL	Biochemical Genetics	257
Ornithine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Osmolality, measured	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Osmolality, urine	Х	CMP, CMP1	Clinical Microscopy	152
		CMQ	QCC, Urinalysis	44

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Osmolality, urine (cont.)		LN6	Urine Chemistry CVL	126
		POC3	POC Urine Dipstick Competency	50
	Х	U	Urine Chemistry-General	68
Osteocalcin		BGS	Bone and Growth	85
Oxalate		KSA	Kidney Stone Risk Assessment	69
Oxazepam		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Oxcarbazepine		ZE	Therapeutic Drug Monitoring, Extended	59
Oxcarbazepine metabolite		ZE	Therapeutic Drug Monitoring, Extended	59
Oxidants, urine		DAI	Urine Drug Adulterant/ Integrity Testing	101
Oxycodone		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Oxyhemoglobin	Х	SO SO	Blood Oximetry	95
		SOQ	QCC, Blood Oximetry	41
Oxymorphone		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
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Analyte/Procedure	LAP	Program	Description	Page
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Pancreatic amylase	Х	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
PAPP-A		FP1B	First Trimester Maternal Screening, Free Beta	88
		FP1T	First Trimester Maternal Screening, Total hCG	88
Parainfluenza virus		ID2	Nucleic Acid Amp, Respiratory	204
	Х	IDPN	Infectious Disease, Pneumonia Panel	211
	Х	IDR	Infectious Disease, Respiratory Panel	210
	X	VR1	Virology Culture	200
	Х	VR2	Viral Antigen Detection by DFA	200
Paraprotein identification	Х	SPE	Protein Electrophoresis	76
Parasite identification	X	BP	Blood Parasite	198
		GIP, GIPN	Gastrointestinal Panel	212- 213
	Х	GIP5	Gastrointestinal Panel, 5 Challenge	212
	X	MVP	Molecular Vaginal Panel	191
	Х	P, P3, P4, P5	Parasitology	197
	ļ.,	PEX	Expanded Parasitology	198
	X	STIM	Sexually Transmitted Infection Detection, Molecular	191
		TMO	Ticks, Mites, and Other Arthropods	198
		TVAG	T. vaginalis, Molecular	197
	X	TVG5	<i>T. vaginali</i> s, Molecular, 5 Challenge	197
	Х	VS	Vaginitis Screen	190
		WID	Worm Identification	198
Parathyroid hormone (PTH)	X	PTH	Parathyroid Hormone	86
		PTHQ	QCC, PTH	40
Parentage/relationship testing	X	PARF	Parentage/Relationship	246
Paroxetine		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Parvimonas micra		JIP	Joint Infection Panel	208
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pCO ₂	Х	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
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		LN13, LN13C	Blood Gas CVL	128
		POC10, POC11	POC Competency Blood Gases	51
PDGFRA	Х	KIT	KIT/PDGFRA	276
	Х	MTP	Multigene Tumor Panel	277
PD-L1	X	PDL1	PD-L1 Immunohistochemistry	299
Pentobarbital		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Peptoniphilus spp.		JIP	Joint Infection Panel	208
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	X	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		GOCB	Gastric Occult Blood	157
		LN13, LN13C	Blood Gas CVL	128
		POC10, POC11	POC Competency Blood Gases	51
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		CMQ	QCC, Urinalysis	44
		DAI	Urine Drug Adulterant/ Integrity Testing	101
	Х	HCC2	Waived Combination	66
		POC3	POC Urine Dipstick Competency	50
		UDC	Forensic Urine Drug Testing, Confirmatory	102
Phencyclidine		DFC	Drug-Facilitated Crime	111

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Phencyclidine (cont.)		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Phenethylamine		FTC	Forensic Toxicology, Criminalistics	107
Pheniramine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Phenobarbital	X	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		LN3	TDM CVL	125
		Т	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Phentermine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Phenylalanine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Phenylephrine		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Phenytoin	X	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
		LN3	TDM CVL	125
		SC0	Serum Carryover	137
		Т	Toxicology	98
		UT	Urine Toxicology	98
Phenytoin, free	X	CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56

Analyte/Procedure	LAP ENR		Description	Page
Phenytoin, free (cont.)		CZQ	QCC, Chemistry and TDM	37
Phosphorus	Х	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Phosphorus, urine		LN6	Urine Chemistry CVL	126
	Х	U	Urine Chemistry-General	68
PIK3CA	Х	MTP	Multigene Tumor Panel	277
Pinworm prep	Х	CMMP	Clinical Microscopy, Misc	153
Pipette calibration, gravimetric		I	Instrumentation	137
Plasma cell myeloma, measurable (minimal) residual disease		FL9	Flow Cytometry Plasma Cell Myeloma Measurable (Minimal) Residual Disease	228
Plasma cell neoplasms		PCNEO	Flow Cytometry, Plasma Cell Neoplasms	228
Plasma hemogloblin		PHG	Plasma Hemoglobin	76
Plasminogen activator inhibitor		CGE/CGEX	Coagulation, Extended	167
Plasminogen activator inhibitor (PAI)-1 (SERPINE1 gene)		MGL1	Molecular Genetics	262- 263
Plasminogen antigen		CGE/CGEX	Coagulation, Extended	167
Plasmodium falciparum antigen	Х	RML5	Rapid Malaria, 5 Challenge	198
		RMAL	Rapid Malaria	198
Platelet aggregation		PF	Platelet Function	171
Platelet antibody detection	Х	PS	Platelet Serology	238
Platelet calculator		TRC	Transfusion-Related Cell Count	237
Platelet count	X	FH1-FH4, FH9-FH10, FH13, FH16-FH17	Hematology Automated Differential	140
		FH3Q, FH4Q, FH9Q, FH13Q	QCC, Automated Hematology Series	43
	Χ	HE	Basic Hematology	140
		LN9	Hematology CVL	127
Platelet count, estimated		EHE1	Expanded Virtual Peripheral Blood Smear	150

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Platelet count (platelet- rich plasma)	Х	TRC	Transfusion-Related Cell Count	237
Platelet crossmatch		PS	Platelet Serology	238
Platelet function		PF1	Platelet Function	171
Platelet mapping		PLTM	Platelet Mapping	174
Plesiomonas shigelloides		GIP	Gastrointestinal Panel	212
	Х	GIP5	Gastrointestinal Panel, 5 Challenge	212
		GIPN	Gastrointestinal Panel, Global	213
PML/RARA		MH02, MH03	Molecular Hematologic Oncology	278
		MRD2	Measurable (Minimal) Residual Disease	279
Pneumocystis detection		PCP1	Pneumocystis jirovecii, Calcofluor White Stain	196
		PCP2	Pneumocystis jirovecii, DFA Stain	196
		PCP4	Pneumocystis jirovecii, GMS Stain	196
PNH immunophenotype		PNH	Paroxysmal Nocturnal Hemoglobinuria, RBC	229
pO ₂	Х	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
		LN13, LN13C	Blood Gas CVL	128
		P0C10, P0C11	POC Competency Blood Gases	51
Porphobilinogen, urine		UPBG	Porphobilinogen, Urine	70
Posaconazole		AFD	Antifungal Drugs Monitoring	109
Post-immunotherapy analysis, flow cytometry		FL6	Post-immunotherapy Flow Analysis	225
Postanalytical DNA sequencing		SEC	DNA Sequencing Count	264
Postvasectomy sperm count, automated		PV1	Postvasectomy Sperm Count	162
Postvasectomy sperm count, manual	Х	PV	Postvasectomy Sperm Count	162
Postvasectomy sperm presence/absence, manual	X	PV	Postvasectomy Sperm Count	162
Potassium	Х	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
		AQQ, AQHQ, AQSQ	QCC, Critical Care Blood Gas Series	42
	Х	C1, C3/C3X, C4, CZ/ CZX/CZ2X	Chemistry and TDM	54-56

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Potassium (cont.)		CZQ	QCC, Chemistry and TDM	37
		FLD2	Body Fluid Chemistry 2	73
		IFS	Interfering Substances	138
		LN13C	Blood Gas CVL	128
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
		POC10, POC11	POC Competency Blood Gases	51
Potassium, urine		LN6	Urine Chemistry CVL	126
	X	U	Urine Chemistry-General	68
Potassium, vitreous fluid		VF	Vitreous Fluid, Postmortem	104
Prader-Willi/Angelman syndrome	Х	MGL1	Molecular Genetics	262- 263
Prealbumin (transthyretin)	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Х	S2, S4	Immunology, Special	217
Predictive markers by immunohistochemistry	X	GHER2	Gastric HER2	297
	Х	HER2	HER2 by Immunohistochemistry	297
	Х	PDL1	PD-L1 Immunohistochemistry	299
		PM1	CD117 by Immunohistochemistry	295
	Х	PM2	ER, PgR by Immunohistochemistry	297
		PM3	CD20 by Immunohistochemistry	298
		PM5	Immunohistochemistry TMA	295
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Pregabalin		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
		ZE	Therapeutic Drug Monitoring, Extended	59
Prekallikrein		CGE/CGEX	Coagulation, Extended	167
Primidone		CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Pro B-natriuretic peptides (See NT-pro B-natriuretic peptides)				
Procainamide		CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Procalcitonin		LN41	Procalcitonin CVL	134
	Х	PCT	Procalcitonin	76
Progesterone		LN8	Reproductive Endocrinology CVL	127
	Х	Y/YY	Sex Hormones	84
Progesterone receptors by immunohistochemistry		PM2	ER, PgR by Immunohistochemistry	297
Prolactin		LN8	Reproductive Endocrinology CVL	127
	Х	Y/YY	Sex Hormones	84
Proline, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Promethazine		DFC	Drug-Facilitated Crime	111
Propoxyphene		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Propranolol		FTC	Forensic Toxicology, Criminalistics	107
		Т	Toxicology	98
		UT	Urine Toxicology	98
Prostate-specific antigen (PSA)		ABS	Accuracy-Based Testosterone, Estradiol	115
	Х	K/KK	Ligand-General	82
		LN23	PSA CVL	130
Prostate-specific antigen, complexed (cPSA)		K/KK	Ligand-General	82
Prostate-specific antigen (PSA), free, measured	Х	K/KK	Ligand-General	82
Prostatic acid phosphatase (PAP)	Х	K/KK	Ligand-General	82
Protein C		CGE/CGEX	Coagulation, Extended	167
		CGS2	Coag Special, Series 2	168
		LN35	Thrombophilia CVL	133
Protein, confirmatory urine		DSC	Dipstick Confirmatory	157

Analyte/Procedure	LAP ENR		Description	Page
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Protein S		CGE/CGEX	Coagulation, Extended	167
		CGS2	Coag Special, Series 2	168
Protein, CSF	Х	M, OLI	CSF Chemistry and Oligoclonal Bands	74
Protein, total	Х	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
		SPE	Lipoprotein and Protein Electrophoresis	76
Protein, urine		ABU	Accuracy-Based Urine	115
	Х	CMP, CMP1	Clinical Microscopy	152
		CMQ	QCC, Urinalysis	44
		DSC	Dipstick Confirmatory	157
	Х	HCC2	Waived Combination	66
		LN6	Urine Chemistry CVL	126
		POC3	POC Urine Dipstick Competency	50
	Х	U	Urine Chemistry-General	68
Proteus spp.	Х	IDPN	Infectious Disease, Pneumonia Panel	211
		JIP	Joint Infection Panel	208
Prothrombin mutation (F2 gene)	X	MGL1	Molecular Genetics	262- 263
	Х	TPM	Thrombophilia Mutations	265
Prothrombin time		APXBN	Anticoagulant Monitoring, Apixaban	170
	Х	CGB	Basic Coagulation	166
	Х	CGL	Coagulation, Limited	166
		CGLQ	QCC, Coagulation, Limited	46
		CGS1	Coag Special, Series 1	168
		CGS4	Coag Special, Series 4	168
		DBGN	Anticoagulant Monitoring, Dabigatran	170
		FNPX	Anticoagulant Monitoring, Fondaparinux	170

Analyte/Procedure	LAP ENR	Program Code	Description	Page
Prothrombin time (cont.)		POC6	POC PT/INR, CoaguChek XS Plus	50
		RVBN	Anticoagulant Monitoring Rivaroxaban	170
	X	WP3, WP4, WP6, WP9	Whole Blood Coagulation	173
Prothrombin time, dilute		CGE/CGEX	Coagulation, Extended	167
Protonitazene		NOB	Novel Opioids and Benzodiazepines	108
Provider-performed microscopy		CMMP	Clinical Microscopy, Misc	153
PRU test		PIA/PIAX	Drug-Specific Platelet Aggregation	173
Pseudocholinesterase	Х	C7	Pseudocholinesterase	77
Pseudoephedrine		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UT	Urine Toxicology	98
Pseudomonas aeruginosa	Х	IDPN	Infectious Disease, Pneumonia Panel	211
		JIP	Joint Infection Panel	208
Quality Management Tools		QP251	Laboratory Staffing Ratios	25
		QPB10, QPB25	Assessment of Consistency of Body Fluid Morphologic Observations	26
		QPC10, QPC25	Assessment of Consistency of Peripheral Blood Morphologic Observations	27
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		QT10	Critical Values Reporting	33
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		QT17	Outpatient Order Entry Errors	34
		QT2	Blood Culture Contamination	30
		QT3	Laboratory Specimen Acceptability	30
		QT4	In-Date Blood Product Wastage	31
		QT7	Satisfaction With Outpatient Specimen Collection	32
		QT8	Stat Test TAT Outliers	32
Quetiapine		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107

Analyte/Procedure	LAP ENR	0	Description	Page
Quetiapine (cont.)		Т	Toxicology	98
		UT	Urine Toxicology	98
Quinidine		CZ/CZX/ CZ2X, Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Quinine		FTC	Forensic Toxicology, Criminalistics	107
Ranitidine		FTC	Forensic Toxicology, Criminalistics	107
Rapamycin (sirolimus)	X	CS	Immunosuppressive Drugs	58
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	X	GIP5	Gastrointestinal Panel, 5 Challenge	212
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		IDME	Meningitis/Encephalitis Panel	209
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	Х	IDPN	Infectious Disease, Pneumonia Panel	211
		JIP	Joint Infection Panel	208
Streptococcus pneumoniae		IDME	Meningitis/Encephalitis Panel	209
	Х	IDM5	Meningitis/Encephalitis Panel	209
	Х	IDPN	Infectious Disease, Pneumonia Panel	211
		JIP	Joint Infection Panel	208
		SBAS	S. pneumoniae Ag Detection	183
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	Х	D1	Group A Streptococcus Culture/Molecular	179
<u> </u>	X	D6	Rapid Group A Strep	182
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	Х	IDPN	Infectious Disease, Pneumonia Panel	211
		JIP	Joint Infection Panel	208
	Х	MC4	Urine Colony Count Combination	180
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15, free (trilodotriyronine)	Х	C1, C3/C3X,	Chemistry and TDM	54-56
	,	CZ/CZX/ CZ2X	onomioury and 12m	0.00
		CZQ	QCC, Chemistry and TDM	37
	Х	K/KK	Ligand-General	82
T3, total (triiodothyronine)		ABTH	Harmonized Thyroid	116
(tinodotnyronine)	X	C1,C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Χ	K/KK	Ligand-General	82
		LN50	Thyroid CVL	136
T3, uptake and related tests	X	C1,C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Χ	K/KK	Ligand-General	82
T4, free (thyroxine)		ABTH	Harmonized Thyroid	116
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Χ	K/KK	Ligand-General	82
T4, total (thyroxine)		ABTH	Harmonized Thyroid	116
	X	C1,C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Χ	K/KK	Ligand-General	82
		LN50	Thyroid CVL	136
Tacrolimus	Х	CS	Immunosuppressive Drugs	58
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Tapentadol		DFC	Drug-Facilitated Crime	111
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Taurine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
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tCO ₂	Х	AQ, AQH, AQIS	Critical Care Blood Gas	92-93
		AQSQ	Quality Cross Check— Critical Care Blood Gas	42
Temazepam		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		OFD	Oral Fluid for Drugs of Abuse	103
		T	Toxicology	98
		UDC	Forensic Urine Drug Testing, Confirmatory	102
		UT	Urine Toxicology	98
Teriflunomide		ZE	Therapeutic Drug Monitoring, Extended	59
Testosterone		ABS	Accuracy-Based Testosterone and Estradiol	115
		LN8	Reproductive Endocrinology CVL	127
	Χ	Y/YY	Sex Hormones	84
Testosterone, bioavailable, measured		Υ	Sex Hormones	84
Testosterone, free, measured		Υ	Sex Hormones	84
Tetrahydrozoline		DFC	Drug-Facilitated Crime	111
Thallium, urine		TMU	Trace Metals, Urine	106
Thallium, whole blood		TMWB	Trace Metals, Whole Blood	106
Theophylline	X	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN3	TDM CVL	125
Threonine, quantitative		BGL2	Amino Acid Quantitation for Inherited Metabolic Disorders	258
Throat culture/molecular	Х	D1	Group A Streptococcus Culture/Molecular	179
	Х	MC4	Urine Colony Count Combination	180
	Х	RMC	Routine Microbiology Combination	180

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Thrombin time		CGE/CGEX	Coagulation, Extended	167
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		DBGN	Dabigatran	170
		ECF	Expanded Coagulation Factors	167
Thrombophilia mutations	X	TPM	Thrombophilia Mutations	265
Thyroglobulin	Х	TM/TMX	Tumor Markers	89
Thyroid-stimulating hormone (TSH)		ABS	Accuracy-Based Testosterone and Estradiol	115
		ABTH	Harmonized Thyroid	116
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Х	K/KK	Ligand-General	82
		LN50	Thyroid CVL	136
Thyroxine (T4), free		ABTH	Harmonized Thyroid	116
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	X	K/KK	Ligand-General	82
Thyroxine (T4), total		ABTH	Harmonized Thyroid	116
	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Х	K/KK	Ligand-General	82
		LN50	Thyroid CVL	136
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Tissue parasite identification	Х	BP	Blood Parasite	198
	Χ	P	Parasitology	197
		PEX	Expanded Parasitology	198
Tobramycin	Х	CZ/CZX/ CZ2X,Z	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		LN3	TDM CVL	125
Topiramate		DFC	Drug-Facilitated Crime	111
		FTC	Forensic Toxicology, Criminalistics	107
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		UT	Urine Toxicology	98
		ZE	Therapeutic Drug Monitoring, Extended	59
Total bile acids		TBLA	Total Bile Acid	77

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Total bilirubin	Х	C1, C3/C3X, C4, CZ/ CZX/CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD2	Body Fluid Chemistry 2	73
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
	X	NB, NB2	Neonatal Bilirubin	64
Total bilirubin, urine	X	CMP, CMP1	Clinical Microscopy	152
		DSC	Dipstick Confirmatory	157
	X	HCC2	Waived Combination	66
Total free fatty acids		FCFS	Fecal Fat	75
Total hCG	Х	FP1T	First Trimester Maternal Screening, Total hCG	88
Total hemolytic complement		CH50	Total Hemolytic Complement	222
Total iron binding capacity, measured	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
Total nitrogen, urine		U	Urine Chemistry–General	68
Total nucleated cells		CBT	Cord Blood Testing	240
		SCP	Stem Cell Processing	240
Total nucleated cells manual differential count (body fluid)		HFC/HFCI	Hemocytometer Fluid Count	158
		VBF	Virtual Body Fluid	154
Total nucleated cells (WBC) automated count (body fluid)		ABF1, ABF2, ABF3	Automated Body Fluid	154
Total protein	X	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
		IFS	Interfering Substances	138
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
		SPE	Protein Electrophoresis	76

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Total protein, CSF	Х	M, OLI	CSF Chemistry and Oligoclonal Bands	74
Total protein, urine	Х	CMP, CMP1	Clinical Microscopy	152
		CMQ	QCC, Urinalysis	44
	Χ	HCC2	Waived Combination	66
		LN6	Urine Chemistry CVL	126
	Χ	U	Urine Chemistry-General	68
Total tricyclics	Χ	SDS	Serum Drug Screen	104
	Х	ZT	TDM, Special	59
Touch imprint/crush prep		TICP, TICP1	Touch Imprint/Crush Prep	309
Toxicology, serum, qualitative	Х	SDS	Serum Drug Screen	104
	Х	T	Toxicology	98
Toxicology, urine, qualitative	X	DMPM	Drug Monitoring for Pain Management	110
	Х	T	Toxicology	98
	Х	UDS, UDS6	Urine Drug Screen	100
	Х	UT	Urine Toxicology	98
Toxicology, urine, qualitative/quantitative	Х	DMPM	Drug Monitoring for Pain Management	110
	Χ	UDC	Forensic Urine Drug Testing, Confirmatory	102
Toxoplasma gondii	X	VR3	Antibody Detection— Infectious Disease Serology	214
TPMT		PGX3	Pharmacogenetics	264
Tramadol		DFC	Drug-Facilitated Crime	111
		DMPM	Drug Monitoring for Pain Management	110
		FTC	Forensic Toxicology, Criminalistics	107
		T	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Transferrin	X	C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54–56
		CZQ	QCC, Chemistry and TDM	37
		LN7	Immunology CVL	126
	Χ	S2, S4	Immunology, Special	217
Transfusion medicine		ETME1	Expanded Transfusion Medicine Exercises	242
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		JATE1	Transfusion Medicine, Automated, Educational	233
		JE1	Transfusion Medicine, Educational	232
		TMCA	Transfusion Medicine, Competency Assessment	239

Analyte/Procedure	LAP ENR		Description	Page
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		TMCAE	Transfusion Medicine, Competency Assessment	239
		TMCAF	Transfusion Medicine, Competency Assessment	239
	Х	TRC	Transfusion-Related Cell Count	237
Trazodone		FTC	Forensic Toxicology, Criminalistics	107
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		UT	Urine Toxicology	98
Treponema pallidum	Х	G	Syphilis Serology	222
Trichomonas vaginalis	Х	MVP	Molecular Vaginal Panel	191
	Х	STIM	Sexually Transmitted Infection Detection, Molecular	191
	X	TVG5	Trichomonas vaginalis, Molecular, 5 Challenge	197
		TVAG	Trichomonas vaginalis, Molecular	197
	Х	VS , VS1	Vaginitis Screen	190
Tricyclic group		Т	Toxicology	98
		UDS, UDS6	Urine Drug Screen	100
		UT	Urine Toxicology	98
Tricyclics, total	X	SDS	Serum Drug Screen	104
	X	ZT	TDM, Special	59
Triglycerides		ABL	Accuracy-Based Lipid	114
	X	C1, C3/C3X, C4, CZ/ CZX/CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
		FCFS	Fecal Fat	75
		FLD	Body Fluid	72
		FLDQ	QCC, Body Fluid Chemistry	38
	X	LCW	Chemistry-Ltd, Waived	64
		LN2	Chemistry, Lipid, Enzyme CVL	124
		LN2BV	Chemistry, Lipid, Enzyme CVL – all Beckman (except AU), Vitros	124
Triiodothyronine (T3), total		ABTH	Harmonized Thyroid	116
	Х	C1, C3/C3X, CZ/CZX/ CZ2X	Chemistry and TDM	54-56
		CZQ	QCC, Chemistry and TDM	37
	Х	K/KK	Ligand-General	82
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POC14	52	RT4*	146	TICP	309	VM1*	243
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POC16	52	RUB*	216	TM*	89	VM3*	243
PS*	238	RUBX*	216	TMB	273	VM4*	244
PTH*	86	RUR*	189	TMCA	239	VM5*	244
PTHQ*	40	RVBN*	170	TMCAD	239	VM6*	244
PV*	162	RWBC*	147	TMCAE	239	VM6X*	244
PV1*	162	S2*	217	TMCAF	239	VPBS*	149
QF*	221	S4*	217	TMO*	198	VR1*	200
QP251	25	S5*	217	TMU*	106	VR2*	200
QPB10	26	SALC*	77	TMWB*	106	VR3*	214
QPB25	26	SARC	275	TMX*	89	VR3M*	214
QPC10	27	SBAS*	183	TPM	265	VR4*	200
QPC25	27	SC*	162	TRC*	237	VRE*	192
QPD10	28	SC1*	162	TTD*	214	VS*	190
QPD25	28	SCDD*	108	TVAG*	197	VS1*	190
QT2	30	SCM1*	159	TVG5*	197	VS2*	192
QT3	30	SCM2*	159	U*	68	WBCR*	66
QT4	31	SCO	137	UAA*	156	WBGQ*	37
QT7	32	SCP*	240	UAA1*	156	WID*	198
QT8	32	SCS*	148	UBJP*	76	WP3*	173
QT10	33	SDS	104	UDC*	102	WP4*	173
QT16	34	SE*	221	UDS*	100	WP6*	173
QT17	34	SEC	264	UDS6*	100	WP9*	173
R*	78	SEC1	264	UDSM	112	WP10*	173
RAG*	235	SFLC*	223	UHCG*	160	γ*	84
RAP*	89	SM*	162	UMC*	160	YBC*	195
RBCAT*	235	SM1CD*	162	UPBG*	70	YVM	90
RDS*	221	SM2CD*	162	URC*	157	YY*	84
RETT	265	SMCD*	162	UT*	98	Z*	54-56
RF*	216	S0*	95	UTCO	137	ZAP70*	230
RFAV1	229	SOQ*	41	UVM	70	ZE*	59
RFAV3	229	SP*	189	V*	223	ZT*	59
RFX*	216	SP1*	189	VBDM*	206		
RHCVW*	244	SPE*	76	VBF*	154	1	
11110444	477	- OI L	7.0	V D1	104	-	

VBP

VBP1

VES*

189

190

79

198

180

198

SPN*

STFR*

ST*

RMAL*

RMC*

RML5*

284

284

172

^{*}Program is ISO/IEC 17043 accredited.

Accreditation to ISO 17043:2010 for proficiency testing

The College of American Pathologists (CAP), the leading organization of board-certified pathologists, serves patients, pathologists, and the public by fostering and advocating excellence in the practice of pathology and laboratory medicine worldwide.

As an accrediting organization ourselves, we recognize the value in having an independent assessment of our management system for our proficiency testing programs. That's why the CAP is accredited by the ANSI National Accreditation Board (ANAB) to the international standard ISO 17043:2010 for proficiency testing.

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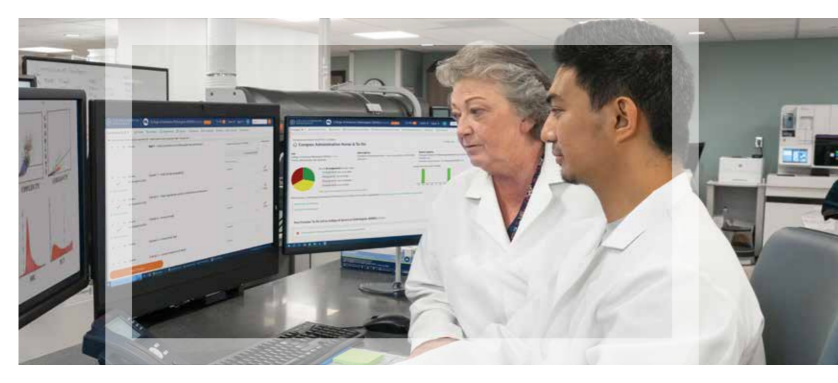
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Notes

Notes

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