

Accredited Medical Test Site (MTS) Application Packet

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Important Information:

Laboratories licensed by the Washington Medical Test Site (MTS) licensure program are exempt from the Clinical Laboratory Improvement Amendments of 1988 (CLIA). You do not need to apply to the Centers for Medicare and Medicaid Services (CMS) for a CLIA number. Your MTS license will contain both your MTS license number and your CLIA number.

Accredited: If you are applying for an accredited license, your MTS must be inspected by the accreditation organization. In facilities, such as hospitals, where testing may be performed at different locations, all areas of laboratory testing must be covered by an MTS license. It is the facility's choice whether to include point of care (ancillary) testing under the same MTS license as the main laboratory, or license separately. Please coordinate with your administration to ensure that all testing is licensed.

Proof of accreditation or certification by the accreditation organization must be included with your application along with documentation for qualifications of the Laboratory Director (such as degrees, board certifications, and CV's).

If your MTS is located in a facility accredited by the Joint Commission, you have the option of being licensed as a Categorized Medical Test Site and inspections will be performed by the Washington State Medical Test Site (MTS) Program. If your facility is currently accredited by the Joint Commission and you choose to have the MTS program do the laboratory inspection, do not complete this application. Complete the Categorized MTS Application Packet.

Per <u>WAC 246-338-050</u>, all licensed medical test sites, excluding those granted a certificate of waiver, must enroll in proficiency testing for all CMS regulated analytes.

In order to process your request:

Mail your application with initial documentation and your check or money order payable to:

Department of Health P.O. Box 1099 Olympia, WA 98507-1099 Contact Us: 360-236-4985

To request this document in another format, call 1-800-525-0127. Deaf or hard of hearing customers, please call 711 (Washington Relay) or email doh.information@doh.wa.gov.

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Fee Information

Initial - Submit the fee corresponding to the license Category your site falls into based on your site's test volume and number of testing specialties.

The categories are based on the number of specialties (SPEC) performed and the estimated annual volume of testing. MTS accredited license applications received during the first year of the state biennium (7/01/2023 through 6/30/2024) are required to submit the full fee. Applications received during the second year of the state biennium (7/01/2024 through 6/30/2025) are required to submit half of the full fee. The license categories and corresponding fees are:

Category	Fee – Applies to applications submitted during the first year of the biennium 7/01/2023 – 6/30/2024	Fee – Applies to applications submitted during the second year of the biennium 7/01/2024 – 6/30/2025
Low Volume (1-2000)	\$ 230	\$115
A (2,001-10,000, 3 SPEC)	\$ 290	\$145
B (2001-10,000, 4 SPEC)	\$ 320	\$160
C (10,001-25,000, 3 SPEC)	\$ 730	\$365
D (10,001-25,000, 4 SPEC)	\$ 780	\$390
E (25,001-50,000)	\$ 1,090	\$545
F (50,001-75,000)	\$ 1,740	\$870
G (75,001-100,000)	\$ 2,390	\$1,195
H (100,001-500,000)	\$ 3,090	\$1,545
I (500,001-1,000,000)	\$ 8,920	\$4,460
J (>1,000,000)	\$ 11,330	\$5,665



Accredited Medical Test Site Application Instructions Checklist

When your application for a Medical Test Site is received by the Department of Health, you will be notified in writing of any outstanding documentation needed to complete the application process.

All information should be printed clearly in blue or black ink. It is your responsibility to submit the required forms.

Indicate type of application:

- New Choose this option if the facility has never been issued an MTS license.
- Change of ownership Choose this option if the facility was previously issued an MTS license and is now under new ownership and/or has a new UBI number.
- Change of license type Choose this option if the facility has previously been issued a different type of MTS license, such as a waived MTS license, a PPMP MTS license, or a categorized MTS license.

☐ Check One:

Please check your legal owner/operator business structure type according to your Washington State Master Business License.

Section 1. Demographic Information:

Unified Business Identifier Number (UBI #): Enter your Washington State UBI #. All Washington State businesses must have a UBI #. City, county, and state government departments also have UBI #s.

Federal Employer ID Number (FEIN): Enter your FEIN, if the business has been issued one. If the facility FEIN # is different than the Legal Owner FEIN, enter this number on page two of the application under Facility Specific Federal Employer ID Number (FEIN).

Legal Owner/Operator Name: Enter the owner's name as it appears on the UBI/Master Business License.

Legal Owner Mailing Address: Enter the owner's complete mailing address.

Phone and Fax: Enter the owner's phone and fax numbers.

Email and Web Address: Enter the owner's email and facility web addresses, if applicable.

Facility Name: Enter the lab's name as advertised on signs and web site.

Facility Specific Federal Employer ID Number (FEIN). Enter if different from the Owner FEIN listed on page one of the application.

Physical Address: Enter the lab's physical street location including city, state, zip code, and county.

Phone and Fax Numbers: Enter the lab's phone and fax number.

Mailing Address: Enter the lab's mailing address, if different than physical address.

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Section 2. Facility Specific Information: Site Type: Please check one applicable site type.
Hours of Laboratory Testing: List the days and hours of testing for this site.
Additional locations under this license: Attach a list of names, addresses and phone numbers for additional locations, if applicable, and test(s) performed at each site.
Section 3. Key Individuals: Lab Director: Enter the lab director's:
 First name, Last name, and Washington State professional license number, if applicable. (See Section 5. Personnel Qualification Requirements)
2. Email address
Lab Contact: Enter the lab contact's:

- 1. First name, Last name, and Washington State professional license number, if applicable.
- 2. Email address

The lab contact will receive all information that we mail to your medical test site.

Section 4. Additional Information:

Waived Tests: Fill in the test system and test manufacturer in the provided table for each test your lab performs. Refer to the CLIA waived test list provided by the FDA to verify the test you are using is approved for waived use.

PPMP Tests: Next to each test, provide an annual estimate of the volume of testing to be performed. The microscopic procedures can only be performed in your facility by a Washington State licensed MD, DO, DPM, ARNP, PA, or dentist.

Non-Waived Tests: Place a checkmark by all the non-waived tests performed at your medical test site. If the tests performed are not listed, add the tests under the appropriate specialty/subspecialty (bold headings). For volumes, provide an estimate of the annual number of tests to be performed. Attach additional sheets if needed. Do not include waived or PPMP tests when counting volumes. Proof of enrollment in proficiency testing must be submitted with the application.

Use the following guidelines for counting tests:

Allergens: count each individual allergen as one test.

Chemistry profiles: count each individual analyte separately.

Complete blood counts: count each measured individual analyte separately that is ordered and reported separately. Differentials are counted as one test. Manual differentials are counted as a separate test.

Cytogenetics: the number of tests is determined by the number of specimen types processed on each patient; e.g., a bone marrow and a venous blood specimen received on one patient is counted as two tests.

Cytology: count each slide (not case) as one test for both pap smears and nongynecologic cytology.

Histocompatibility: count each HLA typing (including disease associated antigens), HLA anti-body screen, or HLA crossmatch as one test.

Histopathology: count each block (not slide) as one test. Autopsy services are not included.

For those laboratories that perform special stains on histology slides, the test volume is determined by adding the number of special stains performed on slides to the total number of specimen blocks prepared by the laboratory.

Immunohematology: count each ABO, Rh, antibody screen, crossmatch, or antibody identification as separate tests.

Microbiology: count susceptibility testing as one test per group of antibiotics used to determine sensitivity for one organism. Count cultures as one per specimen regardless of the extent of identification, number of organisms isolated and number of tests/procedures required for identification.

Urinalysis: count microscopic and macroscopic examinations as separate tests. Count macroscopics (dipsticks) as one test regardless of the number of reagent pads on the strip.

Section 5. Personnel Qualification Requirements: Personnel Qualification Requirements (Moderate & High Complexity Testing): The laboratory director must submit a copy of evidence of qualifying credentials and training.

The following courses are available to obtain 20 CE credit hours in laboratory practice that cover the laboratory director responsibilities:

- University of Iowa CLIA-CME Course for Physician Lab Directors of Moderate Complexity Laboratories: https://cme.medicine.uiowa.edu/
- COLA's Laboratory Director CME Certification Course: https://education.lms.cola.org/catalog/info/id:133
- COLA's Annual Laboratory Enrichment Forum: https://education.cola.org/2024-laboratory-enrichment-forum
- LabUniversity Laboratory Director CME Program: https://labuniversity.org/lab-director-cme-program/

Section 6. Other Licensure, Certification, or Registration Information: Legal Owner: List the names, titles, addresses, and phone numbers of the corporate officers, LLC members or manager, partners, etc. Attach additional pages, if necessary. Indicate if you wish to retain the CLIA number if switching to a new license type.
Change of Ownership Information: If applicable, list the previous legal owner name, previous name of facility, previous MTS license number, effective date of ownership change and physical address. Indicate if you wish to retain the CLIA number if changing ownership.
Section 7. Foreign Ownership: Complete if facility is owned fully or partially by a foreign entity.

Signature:	
The legal owner or authorized representative must sign and date the application.	Print the
name and title of the legal owner or authorized representative.	

You will receive a renewal notice for this license approximately 60 days before the expiration date. The renewal will be mailed to the facility mailing address on file.

Please contact Facilities Customer Service at 360-236-4985 if you have any questions or need assistance in completing the application form. Additional information is available on our website at: http://www.doh.wa.gov/mts.



Proficiency Testing (Not required for Waived or PPMP testing)

Proficiency testing (PT), as required under Medical Test Site <u>WAC 246-338-050</u>, is a source of external quality control. This practice of testing unknown specimens from an outside source provides an additional means to assure quality laboratory testing results. Although laboratories perform daily internal quality control with their test systems, external quality control provides important interlaboratory comparisons to determine the accuracy and reliability of your testing procedures.

Accredited Medical Test Sites must enroll in PT for **all regulated analytes** listed on the next page. Proof of enrollment in PT must be submitted with the application. Most programs are offered as five-sample modules shipped in three separate test events annually. A list of the currently approved PT programs and their phone numbers can also be found on the next page. Call the program or check their website for a free copy of their PT brochure.

You must enroll in programs that cover the testing that you are performing. Generally, most programs are five-sample modules shipped in three test events during the exam. **All regulated analytes** must be covered by PT under the five-sample program.

Information needed to enroll:

- The name of your MTS exactly as it appears on your MTS license,
- Address,
- CLIA ID number, and;
- MTS license number.

Select the appropriate program(s) for your laboratory. When enrolling in the PT program(s), you must indicate that a copy of your PT results be sent to the Washington Medical Test Site Program. **This must be done for each analyte**.

For PPMP procedures and moderate and high complexity tests that are not on the regulated analyte list, you must have a means of establishing the accuracy of the procedure two times a year (biannual verification). Some PT providers offer two-sample programs that can be used for biannual verification of tests that are not included on the regulated analyte list.

What must I do if I add a new test? You must notify our office within 30 days and if this new test is a regulated analyte, you must cover the test in the next PT event. When you notify us, we will remind you to enroll in PT and ask you for proof of enrollment.

What if I decide to stop testing an analyte? You must notify our office within 30 days that you have stopped testing. If you have signed up for PT for this analyte, be sure to notify your PT provider and/or choose the code "test not performed" on the PT answer sheet.

If you have other questions, email MTS@doh.wa.gov for assistance.

Additional information is available at our <u>website</u> in the proficiency testing section.

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Approved Proficiency Testing Providers

Accutest 800-665-2575
Amer. Assoc. of Bioanalysts 800-234-5315
American Proficiency Institute 800-333-0958

College of American Pathologists (CAP) 847-832-7000 WSLH 800-462-5261

Regulated Analytes:

Each laboratory must enroll in PT in the following tests:

Chemistry

ALT/SGPT Albumin

Alkaline phosphatase

Amylase AST/SGOT

B-natriuretic peptide

(BNP)

Bilirubin, total (or neonat.) Blood gas p02, pC02, pH

Calcium, total

Cancer antigen (CA) 125

Carbon dioxide

Carginoembryonic antigen

Chloride

Cholesterol, total

LDL cholesterol, direct

HDL cholesterol Creatine kinase

Creatine kinase isoen-

zymes
Creatinine
Ferritin
GGT
Glucose

Hemoglobin A1c

Iron, total

Total iron binding capacity,

direct LDH

ProBNP

Magnesium Phosphorus Potassium

Prostate specific antigen

Sodium Total protein Triglycerides Troponin I Troponin T Urea nitrogen Uric acid

Endocrinology

Cortisol
Estradiol
Free thyroxine
Folate, serum

FSH

Serum pregnancy (HCG) (qualitative or quantitative)

Luteinizing hormone Parathyroid hormone

Progesterone Prolactin Testosterone T3 uptake

Triiodothyromine TSH -Thyroxine Vitamin B12

Toxicology

Acetaminophen, serum Alcohol, blood Blood lead Carbamazepine Digoxin

Gentamicin
Lithium
Phenobarbital
Phenytoin
Salicylate
Tobramycin
Theophyline
Valproic acid
Vancomycin

Hematology

Cell identification
Auto or manual WBC diff.
Erythrocyte count (RBC)
Hematocrit (automated)
Hemoglobin
Leukocyte count (WBC)
Platelet count
Fibrinogen
Partial thromboplastin

Prothrombin time

Immunohematology

ABO group
D (Rh typing)
Unexpected Antibody
detection
Compatibility testing
Antibody identification

Syphilis Serology

RPR, VDRL, MHA-TP, etc.

Immunology

Alpha–1 antitrypsin AFP (tumor marker) Antinuclear antibody Anti-HCV ASO C-reactive protein (high sensitivity) HIV

Immunology (cont.)

Complement C3, C4

HBsAg, Anti-HBc, HBeAg, Anti-HBs, IgA, IgE, IgG, IgM Infectious mononucleosis Rheumatoid factor Rubella

Bacteriology

Chlamydia
Direct Strep test
GC
Throat culture
Urine culture ID
Gram stain

Other culture/combina-

tions

Antimicrobial tests

Mycology

Yeast ID/culture Fungus culture—systemic

Parasitology

Direct only Concentration/Stain

Virology

HSV EIA Culture or FA Other EIA for virus

Mycobacteriology

AFB Smear and/or culture

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Olympia, WA 98507-1099 360-236-4700 http://www.doh.wa.gov/mts Date Stamp Here

Accredited Medical Test Site License Application				
This is for: New Change of	Ownership	Change of Lie	cense Type	
Check One				
Association L	imited Partne	ership 🔲 Parti	nership	
☐ Corporation ☐ N	/lunicipality (C	City) Sole	Proprietor	
	/lunicipality (C	• /	e Government Agency	
	Ion-Profit Cor	poration Trus	t	
Section 1. Demographic Info				
UBI#	Fe	ederal Employer ID Nu	ımber (FEIN)	
Legal Owner/Operator Entity Name (as it appe	ears on the U	BI/Master Business Li	cense)	
Mailing Address				
City	State	Zip Code	County	
Phone (enter 10 digit #)		Fax (enter 10 digit	#)	
Email Address		Web Address		
Facility/Agency Name (Business name as adv	ertised on sig	gns or website)		
Facility Specific Federal Employer ID Number	(FEIN) (if diff	erent than one entere	d above.)	
Physical Address				
	T			
City	State	Zip Code	County	
Facility Phone (enter 10 digit #)		Facility Fax (enter	10 digit #)	
Mailing Address (If different than physical address)				
City	State	Zip Code	County	
•		·		
For Office Use Only				
Medical Test Site #		_CLIA#		

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Section	n 2. Facil	ity Specif	ic Informa	tion			
1 Am 2 Am 3 An 4 Ass 5 Blo 6 Co 7 Co 8 En 9 Fe 10 H	e (check one or abulance abulatory Surger cillary Test Site sisted Living Fac ood Banks ammunity Clinic amprehensive Ood d Stage Renal I derally Qualified ealth Fair ealth Main. Orga	ry Center cility utpatient Rehab Disease Dialysis I Health Center	13 Hospi 14 Hospi 15 Indep 16 Indus 17 Insura 0 18 ICFM s 19 Mobile 20 Pharn 21 Physic	tal endent Laborato trial ance R e Lab	ory	23 Prison 24 Public He 25 Rural Hea 26 Student H 27 Skilled Nu 28 Tissue Ba 29 Other 30 Drug Trea 31 Clinic	olth Clinic ealth Service orsing Facility nk/Repository
Accredi	tation Organiz	zation					
The following contains information only for those laboratories that choose to be inspected and accredited by a private accrediting organization. Complete the information below to indicate which accrediting organization will inspect your laboratory. A2LA AABB ACHC ASHI CAP COLA The Joint Commission To qualify for this type of license you MUST include proof of accreditation of the laboratory testing at your facility with your application. If you have not yet been inspected by the accrediting organization, include proof of enrollment with your application, and forward the proof of accreditation after the survey has been completed. If you do not wish to be inspected by a laboratory accrediting organization, you will instead need to apply for a categorized medical test site license.							
Hours o	f Laboratory	Testing					
List days	and times durin	g which laborat	t ory testing is p	performed. If te	sting 24/7 chec	k here	
	Sunday	Monday	Tuesday	Wednesday	Thursday	Friday	Saturday
From:							
To:							
Addition	al locations u	nder this lice	nse				
health tes license. This licen If yes: Att and a list CLIA num	alify as a not-forsting (total of 15 ase will have add ach a list of nan of tests performabers of the site y, you must incl	or less waived ditional location nes, addresses ned at each site s that will be co	or moderate co s under one lice and phone nun . If any of the si nsolidated unde	mplexity tests) a ense and the parabers for each s tes already have er this license. If	ragraph above ite that will be ite a MTS license you are not a s	tions, you may a applies: Yes ncluded under oe, include the Metate or local go	□ No □ license, TS and vernment

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Section 3. Key Individua	Section 3. Key Individuals					
Lab Director (include MD, PhD, BS, e	etc.)					
First Name	Last Name WA	State Professional License number				
Email Address						
Does the director of this laboratory ser Washington or another state?	ve as director for any other laboratories Yes No	s that are separately licensed in				
If yes, provide the name of the laborate	ory and CLIA number:					
Lab Contact Person						
First Name	Last Name WA	State Professional License number				
Email Address						
Section 4. Additional Info	ormation—Waived Tests					
Complete the table below for waived te Section 4, if you need assistance comp		to the Application Instructions Checklist,				
Test Name	Test System (e.g. One Step Glucose)	Test Manufacturer (e.g. ACME)				
Adenovirus						
Aerobic/Anaerobic Organisms - Vaginal						
Alanine Aminotransferase (ALT) (SGPT)						
Albumin						
Albumin, Urinary						
Alcohol, Saliva						
Alkaline Phosphatase (ALP)						
Amines						
Amphetamines						
Amylase						
Aspartate Aminotransferase (AST) (SGOT)						
Bacteria Associated With Bacterial Vaginosis						
Barbiturates						
Benzodiazepines						

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Bilirubin, Total

Waived Tests (continued)	
Bladder Tumor Associated Antigen	
B-Type Natriuretic Peptide (BNP)	
Buprenorphine	
Calcium, Ionized	
Calcium, Total	
Cannabinoids (THC)	
Carbon Dioxide, Total (CO2)	
Catalase, Urine	
Chlamydia	
Chloride	
Cholesterol	
Cocaine Metabolites	
Collagen Type I Crosslink, N-Telopeptides (NTX)	
Cotinine	
Creatine Kinase (CK)	
Creatinine	
Eddp (Methadone Metabolite)	
Erythrocyte Sedimentation Rate (ESR), Nonautomated	
Estrone-3 Glucuronide	
Ethanol (Alcohol)	
Fecal Occult Blood	
Fentanyl	
Fern Test, Saliva	
Follicle Stimulating Hormone (FSH)	
Fructosamine	
Gamma Glutamyl Transferase (GGT)	
Gastric Occult Blood	
Gastric pH	
Glucose	
Glycated Hemoglobin, Total	
Glycosylated Hemoglobin (HGB A1C)	
hCG, Urine	
HDL Cholesterol	
Helicobacter Pylori	
Helicobacter Pylori Antibodies	
Hematocrit	
Hemoglobin	
Hemoglobin By Copper Sulfate, Nonautomated	
Hepatitis C Virus Antibody	
Herpes Simplex I And/Or II Antibodies	
HIV-1 And HIV-2 Antibodies	

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Waived Tests (continued)	
HIV-1 And HIV-2 Antigens	
Infectious Mononucleosis Antibodies (Mono)	
Influenza (A/B)	
Ketone, Blood	
Ketone, Urine	
Lactic Acid (Lactate)	
LDL Cholesterol	
Lead, Blood	
Leukocyte Esterase, Urinary	
Lithium	
Luteinizing Hormone (LH)	
Lyme Disease Antibodies (Borrelia Burgdorferi Abs)	
Matrix Metalloproteinases-9 (MMP-9)	
Methadone	
Methadone Metabolite (EDDP)	
Methamphetamine	
Methylenedioxymethamphetamine (MDMA)	
Microalbumin	
Morphine	
Neisseria Gonorrhoeae	
Neutrophil Percentage (Neut%)	
Nicotine And/Or Metabolites	
Nitrite, Urine	
Norfentanyl	
Nortriptyline	
Opiates	
Osmolality, Tears	
Ovulation Test (LH) By Visual Color Comparison	
Oxazepam	
Oxycodone	
рН	
pH, Urine	
Phencyclidine (PCP)	
Phenobarbital	
Phosphorus	
Platelet Aggregation	
Platelet Count	
Potassium	
Pregnanediol Glucuronide	
Propoxyphene	

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Waived Tests (continued)	
Protein, Total	
Prothrombin Time (PT)	
Red Blood Cell Count (Erythrocyte	
Count) (Rbc)	
Respiratory Bacterial Pathogens	
Respiratory Syncytial Virus	
Respiratory Viruses	
SARS-CoV-2	
SARS-CoV-2 And Other Respiratory Viruses	
Secobarbital	
Semen	
Sodium	
Spun Microhematocrit	
Streptococcus, Group A	
Thyroid Stimulating Hormone (TSH)	
Tramadol	
Treponema Pallidum (Syphilis) Antibodies	
Trichomonas	
Tricyclic Antidepressants	
Triglyceride	
Urea (BUN)	
Uric Acid	
Urinary Protein, Qualitative	
Urine Dipstick Or Tablet Analytes, Nonautomated	
Urine hCG By Visual Color Comparison Tests	
Urinalysis	
Vaginal pH	
White Blood Cell Count (Leukocyte Count) (WBC)	
White Blood Cell Differential (WBC Diff)	
Whole Blood Qualitative Dipstick Glucose	
Yeast, Candida Only	
Other Waived Test(S) Not Listed	

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Provider-Performed Microscopic F	rocedures (PPIVIP)			
Next to each microscopic procedure, a state licensed MD, DO, DPM, ARNP, you need assistance completing this tab	PA or dentist. Refer to the Application	-		
Check all that apply Direct wet mount preparations for the proof bacteria, fungi, parasites, and human Fecal leukocyte examinations Fern tests Nasal Smears for granulocytes Pinworm examinations	cellular elements	 Post-coital direct, qualitative examinations of vaginal or cervical mucous Potassium hydroxide (KOH) preparations Qualitative semen analysis (limited to the presence/ absence of sperm and detection of motility) Urine sediment examinations 		
Non-waived and Non-PPMP Tests				
Refer to the Application Instructions Che All analytes listed in bold print are regula submitted with the application.				
	Microbiology			
Microscopic Procedures	Bacterial Toxin Detection	Yeast Culture		
Total Volume:	Blood Culture	Growth/No Growth		
NOTE: If the following microscopic tests are ONLY done by a licensed provider, DO NOT complete this section	Chlamidia CSF Culture	Culture and ID		
Wet Mounts	Gram Stain	Parasitology		
Fecal Leukocytes	GC	Total Volume:		
KOH	Throat Culture	Direct Smear		
Pinworm	Urine Culture	Concentrate/Stain		
Post Coital Vagina Mucous Exam Fern Tests	Urine Colony Count Other Culture/ID:	Parasitic Antigens		
Qualitative Semen Analysis (post		Virology		
vas)	Mycobacteriology	Total Volume:		
Quantitative Semen Analysis	Total Volume:	Herpes Antigen		
Urine Sediment	AFB Smear/Stain	Herpes Culture		
Nasal Smear for Granulocytes	AFB Antibiotic Sensitivities	Other Viral Culture		
	AFB Culture & ID	Viral Antigen Detection		
Bacteriology		Human Papillomavirus (HPV)		
Total Volume:	Mycology	Influenza (nonwaived kits)		
Affirm VP (TV, GV, YST)	Total Volume:	RSV (nonwaived kits)		
Antibiotic Sensitivities	DTM Only	SARS-CoV-2 (nonwaived kits)		
Bacterial Antigens	Direct fungal antigen detection	Other (list):		
Clostridium difficile	Fungus Culture			
Group A Strep) rapid test - nonwaived kits)	Growth/No Growth Culture and ID			
Group B Strep				

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Diagnostic Immunology	Histocompatibility
Syphilis Serology	Histocompatibility
Total Volume:	Total Volume:
RPR	Transplant
VDRL	Nontransplant (list specific tests):
MHA-TP (TP-PA)	
FTA	
General Immunology	Pathology
Total Volume:	Pathology
Allergy Testing (count individual allergens tested)	Total Volume:
Alpha-1 Antitrypsin	Histopathology/year
AFP/Tumor	Dermatopathology/year
AFP/Other	Oral Pathology/year
ANA	Gyn Cytology/year
ASO	Non-gyn Cytology/year
Anti-HCV	
HIV	
C3	
C4	
C-reactive protein	Radiobioassay
C-reactive protein (high sensitivity)	Radiobioassay
HBsAg	Total Volume:
Anti-HBc	
Anti-HBs	(list in vitro tests, i.e. blood volume by Cr 51, Schilling test, etc.)
HBeAg	Do NOT include routine RIA tests
HCV	
lgA	
lgG	
lgE	
IgM	
Infectious Mononucleosis (nonwaived kit)	
Rheumatoid Factor	
H. pylori (nonwaived kits)	
COVID-19 Serology	
Rubella Antibody	
Other (list):	

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Moderate and High Complexity Testing Performed in	your facility (attach additional sheets if needed)				
Che	Chemistry				
Routine Chemistry	Phosphorus				
Total Volume:	Potassium				
NOTE: Each measured parameter must be counted as a	ProBNP				
separate test, added together, and included in the Routine Chemistry total volume above.	PSA (Prostate specific antigen, total)				
ALT/SGPT	Sodium				
Albumin	Total iron binding capacity (TIBC), direct				
Alkaline Phosphatase	Total Protein Triglycerides				
Amylase					
ANT/SGOT	Troponin I				
Bilirubin, Total/Neonatal	Troponin T				
B-natriuretic peptide (BNP)	Urea Nitrogen (BUN)				
pH (blood gas)	Uric Acid				
pri (blood gas)	Ammonia				
pO2 (blood gas)	Bilirubin, direct				
Poz (blood gus) Calcium, Total	C-peptide				
Cancer antigen (CA) 125	CA 19-9				
Carbon Dioxide	CA 15-3				
Carcinoembryonic antigen (CEA)	Ceruloplasmin				
Chloride	FFN (Fetal Fibronectin)				
Cholesterol, Total	Free PSA				
Cholinesterase: RBC methodology:	Haptoglobin				
plasma/serum methodology:	Homocysteine				
HDL Cholesterol	Lactic Acid				
LDL-Direct Cholesterol	Lipase				
Creatine Kinase	Ketones, serum				
CK Isoenzymes	Osmolality				
Creatinine	Protein Electrophoresis				
Ferritin	Prealbumin				
GGT	ROM (Rupture of Membranes)				
Glucose	Transferrin, direct				
Glucose Hemoglobin A1C	Vitamin D				
Iron, Total	Other (list):				
LDH					
LDH Isoenzymes					
Magnesium					
Myoglobin					
, oglobii					

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Chemistry (continued)				
Urinalysis	Toxicology			
Total Volume:	Total Volume:			
Strip by nonwaived instrument	Acetaminophen, serum			
	Alcohol, Blood			
Endocrinology	Carbamazepine			
Total Volume:	Digoxin			
Cortisol	Ethosuximide			
Estradiol	Gentamicin			
Folate, serum	Lead, Blood			
FSH	Lithium			
FT3 (Free Triiodothyronine)	Phenobarbital			
FT4 (Free Thyroxine)	Phenytoin			
HCG (Serum Pregnancy or nonwaived urine HCG)	Primidone			
Luteinizing hormone (LH)	Procainamide/metabolites			
Parathyroid hormone (PTH)	Quinidine			
Progesterone	Salicylate			
Prolactin	Theophylline			
Testosterone	Tobramycin			
T3 Uptake	Valproic Acid			
T3 (Triiodothyronine)	Vancomycin			
TSH	Drugs of Abuse (urine):			
T4 (Thyroxine)	# of Panels X # of Analytes =Total			
Vitamin B12	Fentanyl			
ACTH (Adrenocorticotropic hormone)	Tacrolimus			
DHEA-S	Other (list):			
Insulin				
Procalcitonin				
Other (list):				

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Moderate and High Complexity Testing Performed in your facility (attach additional sheets if needed)						
Hematology		Immunohematology				
Hematology		Immunohematology				
Total Volume:		Total Volume:				
Cell Identification/Manual Differential		ABO Group/year				
		D (Rh) Typing/year				
CBC (Complete Blood Count):	Note: Each measured	Antibody Detection (Screen)/year				
Auto WBC Differential	parameter (automated	Antibody Identification/year				
RBC	differential, RBC,	Compatibility Test (Crossmatch)/year				
Hematocrit	hematocrit(or MCV), hemoglobin, WBC,	Other (list):				
Hemoglobin	platelets) must be					
WBC	counted as a separate test.					
Platelet Count						
		Genetics				
Reticulocyte Count						
Hemoglobin Electrophoresis		Genetic Testing				
Flow Cytometry		Total Volume:				
ESR (Erythrocyte Sedimentation Rate)		Biochemical Genetic Tests (list tests):				
Other (list):						
		Cytogenetic Tests (list tests):				
Coagulation						
Total Volume:		Molecular Genetic Tests (list tests):				
Fibrinogen		NOTE: add LIDV testing under Virgle ru, add Chleraudia and (ar CC				
PTT		NOTE: add HPV testing under Virology, add Chlamydia and/or GC testing under Bacteriology				
Prothrombin Time						
Thrombin Time						
Factor Assays						
Activated Clotting Time						
D-dimer						
Other (list):						

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Section 5. Personnel Qualification Requirements

Confirm the personnel qualification requirements described in this section are met. Per WAC 246-338-060(1)(c), medical test site owners must ensure personnel meet the standards for qualifications and responsibilities in compliance with federal regulation, as listed in 42 C.F.R. Part 493 Subpart M – Personnel for Non-waived Testing. A copy of 42 C.F.R. Part 493 Subpart M will be furnished upon request.

Will the MTS perform high complexity tests?

- If no, review Moderate Complexity Testing Personnel Requirements below.
- If yes, review High Complexity Testing Personnel Requirements below.

Moderate Complexity Testing Personnel Requirements

This section is applicable to medical test sites that perform waived and moderate complexity tests.

Director - Ensure personnel qualifications are met as described in 42 C.F.R. 493.1405. **Submit the following documentation with the application**:

- Diploma or transcripts
- Resume or CV
- · 20 CE credit hours in laboratory practice
- Board certification (if applicable)

Technical Consultant – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1411.

Clinical Consultant – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1417.

Testing Personnel – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1423.

High Complexity Testing Personnel Requirements

This section is applicable to medical test sites that perform waived, moderate and high complexity tests.

Director – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1443. **Submit the following documentation with the application:**

- Diploma or transcripts
- Resume or CV
- 20 CE Credit hours in laboratory practice
- Board certification (if applicable)

Technical Supervisor – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1449.

Clinical Consultant – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1455.

General Supervisor – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1461.

Cytology General Supervisor – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1469, if the subspecialty cytology is performed.

Cytotechnologist – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1483, if the subspecialty cytology is performed.

Testing Personnel – Ensure personnel qualifications are met as described in 42 C.F.R. 493.1489.

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Section 6. Other Li	censure,	Certification,	or R	egistra	tion Infori	nation
Legal Owner Information-	-attach add	itional sheets as ne	eded			
List names, addresses, pho	one numbers,	and titles of corporate	officers	s, partners	, members, ma	nagers, etc.
Name	Address		Phone	e #	Title	
If changing license type, do you	ı want the fac	cility to keep the already	/ assigi	ned CLIA r	number? 🔲 Y	es 🔲 No
If yes, provide the CLIA numbe	r:				-	
Change of Ownership Info	ormation					
Previous Name of Legal Own	er					
Previous Name of Facility		Previous MTS License	e #		Effective Date of Owners	
					Change	
Di : IAII						
Physical Address						
City		State		Zip Code		
Oity		State		Zip Code		
If changing ownership, do you	want the facil	ity to keep the already	assigne	ed CLIA nu	ımber? 🗌 Ye	s 🗌 No
If yes, provide the CLIA number	r:				_	
Section 7. Foreign (lwnorchi	in .				
Section 7. Foreign C)WIIEI SIII	ip				
Does this facility have partial or			_	-		☐ No
If yes, what is the country of ori	gin for the for	reign entity?				
		Signature				
		Signature	-			
			1	-4-4-		41.: 1:
I certify that I have received, recategory. I also certify that the						
3 , ,				,	3	
Cianatura of Our on/Authorizas	d Danrasanta	tive of Madical Toot Cite	_	Dete		
Signature of Owner/Authorized	ı Representa	live of Medical Test Site	3	Date		
			_			
Print Name				Print Title	Э	

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