

Molecular Otolaryngology & Renal Research Laboratories For test inquiries please call: 319-335-6623 • For billing inquiries call: 319-467-1647

Fax: 319-353-5869 or 319-335-9890

Email: morl@uiowa.edu • https://morl.lab.uiowa.edu

CLIA: 16D0966193

MORL - Kidney Testing Requisition Form

REFERRING LABORATORY USE ONLY : please complete below section					F	OR MORL USE ONLY	<i>(</i> :				
Requisition Date:	Comp	leted by:		Accn#:			P	MORL Case #:			
Collection: Blood Da	te/Time:	#Tubes	:Serum	Date/Time:	: <u></u>	#Tubes:	Plasr	na Date/Time:	:	#Tubes:	
Part A) Patient	Information o	r ID Sticke	er <u>(Required)</u>		Part A	1) Patient Demo	graph	nic Information	(Required	<u>D</u>	
Name:					Ethnicit	t y : 🗌 Hispani	с	🗌 Not Hisp	anic		
Last		First			Rad	e : 🗌 White		Black or	African An	nerican	
DOB: /	day /year	_ Sex: (please of	□ Male □ Fe <u> check sex assigned</u>	male d at birth)		Asian		America	n Indian/A	laska Na	ative
Height:		BMI	:			Native H	lawaiia	an or Other Pacif	ic Islander		
						More Th	ian On	e Race			
MRN: Part B) Reporting Information (<u>Required</u>)				Part C) Payment Information (<u>Required</u>) Institutional billing or payment by Visa or MasterCard is accepted.							
Health Care Prov	ider:				***The	MORL will <u>NOT</u> s	submi	t to insurance.			
E-mail Address:					Billing Contact:						
Institution:					Institut	ion:					
Street Address:					Street Address:						
City:	State:		Zip:		City:		State	2:	Zip:		
Phone: ()		FAX: ()		Phone:	()		FAX: ()			
lf	you or your pati	ent would	like to pay by	<mark>/ credit c</mark> a	ard, plea	se contact us at <u>n</u>	norl@	<u>uiowa.edu</u> for in	structions		
Part D) Pertine	<mark>nt Clinical Info</mark>	<mark>rmation (</mark>	<u>Required) – C</u>	omplete	the sec	tion below					
Diagnosis:	aHUS: Trigge	r? 🗌 No [Yes (if <u>yes</u> , de	escribe trigg	ger, eg. BM	T, pregnancy, pneumo	ococcal)	:			
		BGN 🗌 F	PIGN 🗌 TTP	P ST	EC-HUS	Other (comp	olete):				
Family history of renal disease? No Yes (if <u>ves</u> , please provide details in comment & attach a pedigree if available)											
Disease History	Date of sy	mptom or	nset:			Specimen Inform	ation:				
Renal Biopsy:	Yes No	Date:				Was specimen dra	wn	Procedure dat		Deat	NI / A
Hematuria:	Yes No					pre or post:		Procedure dat	e: Pre	Post	N/A
Diarrhea:	Yes 🗌 No					Eculizumab:					
Schistocytes:						PLEX (*affects serol	logies):				
Current Lab Valu	<u>es</u> <u>Value</u>	No	rmal Range	<u>Test [</u>	Data	Renal Tx:					
Hg/Hct:	value		indi Kange	Test	Jale						
Haptoglobin:						BMT (*affects gene	tics):				
Platelets:						Liver Tx:					
sCr/BUN:											
eGFR:						Comments:					
LDH:											
uProt/uCr:											
Urine Blood:											
C3 Level:											
C4 Level:											
ADAMTS13:											



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Patient Name: DOB	8: MRN:			
Please see page 3 for sample handlin	ng requirements - No Weekend Deliveries			
Functional Testing Panels Requested	Genetic Testing Requested			
 C3 Glomerulopathy Complement Panel (C3G-CP) (Serologies for complement-mediated renal diseases) (CH50, APFA, C3b Deposition Assay, FHAA, FBAA, Fluid Phase Activity Assay-IFE, Nephritic Factors (C3Nef-C3CSA, C5Nef-C3CSAP, C4Nef), C3, C3c, C4, FB, Ba, Bb, FD, Properdin, C5, Soluble C5b-9, FH and FI levels) 2 mL frozen serum 2 mL frozen EDTA plasma 	 Genetic Renal Panel (DNA test for TTP, aHUS, HUS, DDD, C3G and other complement diseases) (<i>CFH, CFI, MCP, CFB, CFHR5, C3, THBD, ADAMTS13, PLG, DGKE, G6PD, MMACHC, WT1</i> and MLPA) 3-5 cc EDTA whole blood (room temp or refrigerated) Saliva (OraGene OGR-500 kit) or Buccal swabs, no less than 6 swabs (OraCollect OCD-100) OR 5 µg DNA, minimum concentration 50 ng/µl			
 aHUS (complement-mediated TMA) Panel (aHUS-FP) (CH50, APFA, C3b Deposition, FHAA, FBAA, Fluid Phase Activity-IFE, C3, C3c, C4, FB, Ba, Bb, FD, Properdin, C5, Soluble C5b-9, FH and FI levels) 2 mL frozen serum 2 mL frozen EDTA plasma 	 MLPA Testing ONLY (screening for copy number variations in the CFH-CFHR5 genomic region) 3-5 cc EDTA whole blood (room temp or refrigerated) Saliva (OraGene OGR-500 kit) or Buccal swabs, no less than 4 swabs (OraCollect OCD-100) OR 			
 Autoantibody Panel (AAP) (FHAA, FBAA, Fluid Phase Activity-IFE, Nephritic Factors (C3Nef-C3CSA, C5Nef-C3CSAP, C4Nef) 2 mL frozen serum 	 5 μg DNA, minimum concentration 50 ng/μl Familial Testing (site specific analysis to screen for variants previously identified in a family member) Familial Testing Details: 			
 Complement Biomarker Panel (CBP) (C3*, C3c, C4*, FB, Ba, Bb, C5, Properdin, Soluble C5b-9, FH and FI levels) 2 mL frozen serum* 2 mL frozen EDTA plasma Complement Pathway Activity Panel (CPAP) (CH50, APFA, C3b Deposition Assay) 2 mL frozen serum 	Gene/s: MORL ID# or Variant/s: Relationship to previously tested person: If you are interested in ordering Custom Testing please contact Amy Weaver at 319-335-6623 or <u>amy-weaver@uiowa.edu</u>			
a La Carte Testing Requested	Important Information for <u>ALL</u> Requests			
Autoantibody Tests – 1 mL Frozen Serum FH autoantibody (FHAA) FB autoantibody (FBAA) Fluid Phase Activity (IFE) C3Nef (C3CSA) C5Nef (C3CSAP) C4Nef Biomarker Tests – 1 mL Frozen EDTA Plasma C3 Level (1 mL frozen serum) C3c Level C4 Level (1 mL frozen serum) FD Level	All serum and plasma samples MUST be processed and frozen down to -80° C immediately after collection (please see instructions on page 3). Sample <u>type must be clearly labeled (either serum or plasma)</u> and shipped out <u>overnight</u> on at least 5 lb dry ice (Monday – Thursday). If samples arrive thawed they will be <u>REJECTED</u> . <u>No Weekend Deliveries</u>			
FB Level Ba Level Bb Level Properdin Level C5 Level Soluble C5b-9 Fl Level FH Level Complement Pathway Function Tests – 1 mL Frozen Serum CH50 C3b Deposition Assay APFA (Alternative Pathway Functional Assay)	Molecular Otolaryngology & Renal Research Laboratories For test inquiries please call: 319-335-6623 • Fax: 319-353-5869 For billing inquiries please call: 319-335-6653 • Fax: 319-353-5869 Email: morl@uiowa.edu_ • https://morl.lab.uiowa.edu Ship to: Dr. Richard Smith Molecular Otolaryngology & Renal Research Laboratories The University of Iowa 285 Newton Rd., 5270 CBRB Iowa City, IA 52242-1078 Monday — Friday ONLY – <u>No Weekend Deliveries</u>			
ADAMTS-13 Tests – 1 mL Frozen Citrate Plasma ADAMTS-13 Activity ADAMTS-13 Activity with reflex to Inhibitor (when activity <25%)				



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Molecular Otolaryngology & Renal Research Laboratories Sample Requirements Questions? Contact Amy Weaver at 319-335-6623 or amy-weaver@uiowa.edu

	Questions? Contact Arry weaver at 519-5	35-0023 01	any-weaver@ulowa.eau		
Genetic	Genetic Renal Panel/MLPA/DNA Testing Sample Requirements:				
3-5 cc. EDTA whole blood – <i>MORL not responsible for broken tubes</i>					
•					
٠	5 μg DNA, minimum concentration 50ng/μl (A260/A280 1.8-2) resuspended in 0.1mM EDTA (10mM Tris HCl, 0.1mM EDTA, pH 8,				
	Teknova Cat# T0220)				
•	*Please note: blood samples drawn from a bone marrow transpl	ant patient	will result in genetic results for the <i>donor</i> rather than the		
	patient, buccal swab samples are recommended.				
•	Overnight delivery, Room temperature or refrigerated (DO NOT FREEZE WHOLE BLOOD)				
•	Samples are accepted Monday-Friday.				
•	 Samples may be refrigerated if delivery is delayed (stability – 1 week) 				
Serum (FH & FB autoantibody, APFA, C3b Deposition Assay, CH50, Fluid Phase Activity, C3Nef, C5Nef, C4Nef, C3 and C4 levels) Collection					
Protoco	I (minimum volume 2ml) *PLEX treatments will affect serum tests	s, please w	ait ~14 days after PLEX to draw samples:		
1.	Follow standard phlebotomy techniques to collect at least 6 cc	of whole b	lood drawn in a red-top vacutainer tube.		
	Note: Serum separators with "clot activators" should <u>not</u> be u				
2.	Allow the blood in the red-top tube to clot at room temperatur	-			
3.	Centrifuge the clotted blood at room temperature (1000 x g for	r 10 minute	es).		
4.	Label "Serum" or "Red-top" on clean screw top-tube (s).				
5.	Pipette cell-free supernatant (at least 2 mL) to each labeled tu	be (s).			
6.	Place the tube immediately at -80°C (or on dry ice). Sample <u>m</u>				
	Note: Do not transfer cells with serum. If necessary, centrifug	e a secona	time.		
Plasma	(soluble C5b-9, C3c, Ba, Bb, FD, Properdin, C5, FB, FH, FI levels)	Collection	Protocol (minimum volume: 2ml)		
	eatments will affect plasma tests, please wait ~14 days after PLEX				
1.	Follow standard phlebotomy techniques to collect at least 6 cc				
2.	Centrifuge at room temperature immediate after blood draw (2				
3.	Label "Plasma" or "Lavender-top" on clean screw top-tube(s).	1000 x 8 10	10 minutesj.		
4.	Pipette <u>cell-free supernatant</u> (at least 2 mL) to each labeled tul	be (s).			
5.	Place the tube immediately at -80°C (or on dry ice). Sample m		n deep frozen.		
	Note: Do not transfer cells with plasma. If necessary, centrifu	ge a secon	d time.		
Plasma	(ADAMTS-13 Activity/Inhibitor) Collection Protocol (minimum v	volume: 0	5ml):		
			·		
1.	Follow standard phlebotomy procedure to collect blood in buffered sodium citrate (light blue-top, 3.2%) plastic tubes	1.	Following centrifugation, transfer the top two-thirds of the plasma layer into a new plastic tube.		
	(available in 4.5 mL, 2.7 mL or 1.8 mL full draw tubes).	_			
2.	After collection, invert the tube gently 5 to 6 times.	2.	Re-centrifuge the collected plasma at 1500 to 1800 x g		
3.	Label "Citrate Plasma" or "Blue-top" on clean cryovial screw-		with the <i>brake off</i> for an additional 15-20 minutes to		
•••	top tubes.	2	remove any red cells or platelets.		
4.					
	centrifugation. Samples should be centrifuged between 15		cells at the bottom of the tube.		
	to 60 minutes after blood collection for best results.	7.	Place the tube immediately at -80°C (or on dry ice).		
5.	Re-mix the blood sample immediately prior to centrifugation		Sample <u>must</u> remain deep frozen.		
	by gently inverting the tube 5 to 6 times.				
6.	Centrifuge blood sample at room temperature in a horizontal		Note: if the sample arrives at room temperature a new		
	rotor (swinging bucket rotor) for 15-20 minutes at 1500 to		sample will be required.		
	1800 x g with the <i>brake off</i> .				
Sorum	& Plasma Shipping Requirements:		Ship all samples to:		
•	Serum and plasma must be <u>frozen</u> and shipped OVERNIGHT w	vith a	Dr. Richard Smith		
•	minimum of 3 kg (or 6 lbs) of dry ice.		Molecular Otolaryngology & Renal Research		
•	Cryovials should be put in zip lock bags and completely covere	d in dry	Laboratories		
•	ice to keep the sample frozen until it arrives in the lab.	The University of Iowa			
-		285 Newton Rd., 5270 CBRB			
•	Delivery: Monday-Friday. NO WEEKEND DELIVERIES	Iowa City, IA 52242-1078			
•	Thawed OR unlabeled samples will be REJECTED for testing.		Phone: 319-335-6623		



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Complement Panel tests offered by the MORL:	Test Code:
C3 Glomerulopathy Complement Panel (serologies for DDD, C3GN), Serum and Plasma - CH50, APFA, C3b Deposition Assay, FHAA, FBAA, Fluid Phase Activity Assay-IFE, Nephritic Factors (C3Nef-C3CSA, C5Nef-C3CSAP, C4Nef), C3, C3c, C4, FB, Ba, Bb, FD, Properdin, C5, Soluble C5b-9, FH and FI levels	C3G-CP
aHUS (complement-mediated TMA Functional Panel (serologies for TTP, aHUS, HUS), Serum and Plasma - CH50, APFA, C3b Deposition, FHAA, FBAA, Fluid Phase Activity-IFE, C3, C3c, C4, FB, Ba, Bb, FD, Properdin, C5, Soluble C5b-9, FH and FI levels	aHUS-FP
Autoantibody Panel, Serum - FHAA, FBAA, Fluid Phase Activity-IFE, Nephritic Factors (C3Nef-C3CSA, C5Nef- C3CSAP, C4Nef)	AAP
Complement Biomarker Panel, Plasma - C3, C3c, C4, FB, Ba, Bb, FD, C5, Properdin levels, soluble C5b-9, FH and FI levels	СВР
Complement Pathway Activity Panel, Serum - CH50, APFA, C3b Deposition Assay	СРАР
Autoantibodies to Complement Components	Test Code:
Fluid Phase Activity Assay, Serum (IFE)	07FPA
FH Autoantibody, Serum (ELISA)	07FHAA
FB autoantibody, Serum (ELISA)	07FBAA
C3Nef, Serum (Hemolytic)	06C3NEF
C5Nef, Serum (Hemolytic)	06C5NEF
C4Nef, Serum (Hemolytic)	06C4NEF
Functional Assays of Complement Activity - Pathways	Test Code:
CH50, Serum (Liposome-based method)	07CH50
Alternative Pathway Functional Assay (APFA), Serum (ELISA)	06APFA
C3b Deposition Assay (Hemolytic)	01C3BDA
Complement Protein Biomarkers (including split products)	Test Code:
C3 Level, Serum (Turbidmetry)	07C3L
C3c Level, Plasma (ELISA)	06C3CL
C4 Level, Serum (Turbidmetry)	07C4L
FB Level, Plasma (ELISA)	07FBL
Ba Level, Plasma (ELISA)	06BAL
Bb Level, Plasma (ELISA)	06BBL
FD Level, Plasma (ELISA)	06C5L
Properdin Level, Plasma (ELISA)	06PL
C5 Level, Plasma (ELISA)	06C5L
Soluble C5b-9, Plasma (ELISA)	06SMAC
FI Level, Plasma (ELISA)	07FIL
FH Level, Plasma (ELISA)	06FHL
ADAMTS-13	Test Code:
ADAMTS-13 Activity (a la carte only), Citrate Plasma (FRET)	01ATS13
ADAMTS-13 Activity with reflex to Inhibitor Assay (if activity is <25%), Citrate Plasma (FRET)	01ATS13RFX
Genetic Tests Offered by the MORL:	Test Code:
Genetic Renal Panel: NGS + MLPA (CNVs) for Complement-Mediated Kidney Disease	GRP08
MLPA (CFH-CFHR5): Multiplex Ligation Dependent Probe Amplification	MLPA02

MORL – Kidney Testing Requisition Form - Please see page 2 for Tests Offered

Specimen and shipping requirements along with CPT codes and prices can be found on our website: https://morl.lab.uiowa.edu.



DISCLAIMER:

This request to order molecular diagnostic tests from the MORL certifies to the MORL that the ordering healthcare provider has obtained informed consent from the patient as required by applicable state or federal laws for each test ordered, that the ordering healthcare provider has authorization from the patient permitting the MORL to report results for each test ordered to the ordering physician, and that the ordering healthcare provider assumes responsibility for providing the patient with all associated guidance and counseling regarding the test results.

ALL requested information must be provided, or testing will not be performed

Patient information:	Patient date of birth and gender			
	Patient ethnicity and race			
	Patient's clinical information and family history of kidney disease			

We request extensive patient demographic and clinical information. This information is <u>required</u> as it is very valuable in the interpretation of your patient's results.

Specimen information:	Patient identifiers (full name, date of birth, sex, and medical record number) Date of collection <u>Sample type – frozen samples must be CLEARLY LABELED as either serum or plasma (and type, EDTA or Citrate)</u> Ordering physician
Billing information:	We will <u>NOT</u> bill insurance, Medicare, or patient directly. Institutional billing accepted. Visa and Master Card accepted. Personal checks <u>NOT</u> accepted. Please include contact information including phone & fax number for billing questions.
Reporting Information:	Because of confidentiality issues, reports will only be released to the individual indicated on the page 1 of the testing requisition form.
Research Participation:	If your patient's genetic and functional testing results are inconclusive, they may qualify for research studies on complement-mediated renal diseases that are ongoing at the MORL. If you would like your patient to be considered for this opportunity, please contact Amy Weaver at <u>amy-weaver@uiowa.edu</u> .

IMPORTANT INFORMATION FOR PHYSICIAN OR GENETIC COUNSELOR:

DNA tests may detect an abnormality. Detection methods are greater than 99% accurate. Many of these tests are relatively new. The analysis and interpretation represent our best knowledge and understanding of the genetics of these diseases.

There is a small possibility that a test may not work properly, or an error may occur. You may be asked for an additional sample if it is felt that confirmatory testing is needed.

An error in diagnosis may occur if incorrect information is provided with the sample.

Kidney diseases are complex disorders and penetrance of a phenotype (the degree of kidney disease, for example) may be variable. Research to determine whether a genotype-phenotype correlation exists is ongoing.

Because of the complexity of DNA testing, results should be discussed with a genetic counselor or physician.

Note: Kidney diseases are very complex disorders. This complexity means that variants in many different genes can lead to kidney disease. It is possible that no variants will be detected in the variant screens (the genes) you have requested.