720 Twentieth Street South, Suite 330 Birmingham, Alabama 35294-0005 Phone: (205) 934-5562 Fax: (205) 996-2929

www.uab.edu/medicine/genetics/medical-genomics-laboratory

UAB MGL Accession

For MGL Lab Use Only

	Received:	Reviewed:	Accession:	Billing:	Other:
Initials:					
Date:					
Comment:					

Important Notes

-This form must accompany all specimens received -Billing information (page 5) must be included Additional information is available online at www.ua			-All specimens received must include <u>two</u> patient identifiers and <u>collection date</u> -Testing must be ordered by a qualified clinician					
	Additi	onal information		isition Form		edicai-ge	enomics-iaborat	cory
	Patier	nt Informatio	·			Ord	ering Physic	ian:
Sample Collected: (MM/E				□ Please chec	k box if phy		ould receive rep	
				Name: NPI:				NPI:
Legal Name: (First)		(MI)	(Last)					
				Institution:				
DOB: (MM/DD/YY)		MRN:		Address:				
Address:				City:			State:	Zip:
City:		State:	Zip:	Country:		Phone		
Sex at Birth:		SSN:		Please check presult delivery		□ Fax:		
Parent or Guardian name	(if minor)	:		□ Email:				
	Referrir	ng Lab/Hospit	tal:			Addit	ional Report	ts to:
□ Please check box if lab/	hospital sh	ould receive rep	ort directly	Name:				
Name:				Institution:				
Institution:				Address:				
Address:				City:		State:	Zip:	
City:		State:	Zip:	Country:		Phone	<u> </u> :	
Country:	Phone:			Please check preferred ☐ Fax:				
Please check preferred result delivery:	□ Fax:			result delivery:				
□ Email:								
			Previous Te	esting Histor	γ			
Charle III that and he	☐ Patie	ent or family m	ember is pregnant. LMP:	☐ Patient has had chemotherapy in the past 6 months				rapy in the past 6 months
Check all that apply: Patient has had a bone marrow transplant			☐ Infectious diseases (AIDS, Hepatitis, etc.)					
		На	as this patient or relatives ha	ad previous test	ting? □ Ye	s 🗆 No		
Name/Relationship to patient:			Test/Variant/Lab:					
Name/Relationship to patient:			Test/Variant/Lab:					
			Informe	d Consent				
	test(s) req	uested with the	enefits, limitations, and implica patient/guardian and I have ar e maintained.	ations of genetic	_			

MEDICAL GENOMICS LABORATORY

720 Twentieth Street South, Suite 330 Birmingham, Alabama 35294-0005

Phone: (205) 934-5562 Fax: (205) 996-2929 UAB MGI Accession

Name: DOB: (MM/DD/YY)

Lymphocyte/White Blood Cell-based Comprehensive Testing via Next-Gen Sequencing ☐ RUSH Analysis: Testing completed within 15 working days of receipt of sample (Additional \$600 RUSH fee applied; only available for tests on this page) NF1/Legius syndrome and Other RASopathy Related Conditions NF2/Schwannomatosis/Meningiomatosis ☐ NF1-NG: NGS and Del/Dup: NF1 only ☐ NF2-NG: NGS and Del/Dup: NF2 only ☐ NFSP-NG: NGS and Del/Dup: NF1 and SPRED1 ☐ SCH-NG: NGS: 3 genes: LZTR1, NF2, and SMARCB1; and Del/Dup: □ NNP-NG: NGS: 17 genes (no NF1): BRAF, CBL, HRAS, KRAS, LZTR1, NF2, LZTR1, and SMARCB1 MAP2K1, MAP2K2, NRAS, PPP1CB, PTPN11, RAF1, RASA2, RIT1, SHOC2, SOS1, SOS2, and SPRED1; and Del/Dup: SPRED1 and LZTR1 ☐ MEN-NG: NGS: 4 genes: NF2, SMARCB1, SMARCE1, and SUFU; and Del/Dup: NF2 and SMARCB1 ☐ RAS-NG: NGS: 18 genes: BRAF, CBL, HRAS, KRAS, LZTR1, MAP2K1, MAP2K2. NF1. NRAS. PPP1CB. PTPN11. RAF1. RASA2. RIT1. SHOC2. SOS1. Peripheral Nerve Sheath Tumor Testing SOS2, and SPRED1; and Del/Dup: NF1, SPRED1, and LZTR1 ☐ PNT-NG: NGS: 6 genes: NF1, NF2, KRAS, LZTR1, PTPN11 and ☐ CST-NG: NGS: HRAS only SMARCB1; and Del/Dup: NF1, NF2, LZTR1, and SMARCB1 McCune-Albright Syndrome **Rhabdoid Tumor Predisposition Syndrome** ☐ GNAS-NG: NGS: GNAS exons 8 and 9 only ☐ RT-NG: NGS: SMARCB1 and SMARCA4; and Del/Dup: SMARCB1 only **Tuberous Sclerosis Complex** Capillary Malformation Arteriovenous Malformation Syndrome ☐ TSCP-NG: NGS and Del/Dup: TSC1 and TSC2 ☐ RASA-NG: NGS: and Del/Dup: RASA1 and EPHB4

Additional Information

Test Description Key:

Next Generation Sequencing (NGS) Deletion/Duplication analysis (Del/Dup) For additional testing options via tumor/biopsy, please see page 3 of this order form.

Please contact the lab via phone (205) 934-5562 or via email at medgenomics@uabmc.edu if you have any questions when completing this form. For additional information, visit our website at www.uab.edu/medicine/genetics/medical-genomics-laboratory

Important points of consideration for testing

The average coverage for all of our panels is >1600x. Specifically for the NF1 gene, the NGS approach covers >98% of the NF1 coding region at \geq 350X and 99% ≥200X, allowing detection of very low level mosaicism, down to 3-5% variant allele fraction respectively. For all other genes on our panels, the NGS approach covers an average of 99% at ≥200X. Remaining regions are covered at ≥30X. However, for patients with segmental/mosaic presentation, deep coverage in lymphocyte cells may be insufficient to identify the underlying gene change. Testing the affected tissue(s) may be necessary to confirm a diagnosis. Please see page 3 for our tumor/biopsy-based testing options.

Please note: For patients with an ongoing pregnancy who require comprehensive NF1 testing, "NF1-R" is recommended due to the sensitivity and fast turnaround time of this test (please see page 4 for this option).

Specimen Requirements					
Accepted Specimens	Specimen Information:				
Specimen requirements vary based on test requested; please see our website for more details.	□ Peripheral Blood (EDTA); # Tubes: □ Extracted DNA; Source:				
-Blood: 3-6ml EDTA (receipt within one week of collection) -Saliva: OGR-575 DNA Genotek (kits are provided upon request) -DNA: extracted from lymphocyte cells, a minimum of 25ul at 3µg, O.D. value at 260:280nm ≥1.6 (must be extracted in a CLIA or equivalent certified lab) -Fibroblast cells	□ Saliva (kit must be provided by MGL) □ Other, please describe: Please note: failure to provide a date of collection can delay release of results Sample Collected Date (required):				

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UAB MGL Accession

Name: (First) (MI)) (Last)		DOB: (MM/DD/YY)	
	Tumor/Biopsy-based C	Comprehens	ive Testing	
☐ Please check h	nere if blood or DNA is provided fo	or confirmation	testing. Blood Collected: (MM/DD/YY)	
NF1/SPRED1 on biopsied CAL	s and Neurofibromas	NF2-,	LZTR1-, SMARCB1-related Schwannomatosis	
Please contact the laboratory <u>at least</u> biopsy before ordering this test as media and special shipping instructions apply. B hours of collection	must be provided in advance	☐ NF2-NG: Fresh/Frozen Tumor or Tumor Block for NGS with reflex to Sanger as needed and Del/Dup: <i>NF2</i> only ☐ SCH-NG: Fresh/Frozen Tumor or Tumor Block for NGS with reflex to Sanger as needed and Del/Dup: <i>NF2</i> , <i>LZTR1</i> , and <i>SMARCB1</i>		
☐ NF14C: Sanger(RNA) and Del/Dup: NF1	(with automatic reflex to	R	habdoid Tumor Predisposition Syndrome	
SPRED1) on biopsied CALs NF14N:Sanger(RNA) and Del/Dup: NF1		☐ RT-NG: Fresh/Frozen Tumor or Tumor Block for NGS with reflex to Sanger as needed <i>SMARCB1</i> and <i>SMARCA4</i> ; and Del/Dup: <i>SMARCB1</i> only		
RASopathy Related	Conditions	Meningiomatosis		
□ NNP-NG: Fresh/Frozen Tumor for NG:	` ′		esh/Frozen Tumor or Tumor Block for NGSwith reflex to Sanger as SMARCB1, SMARCE1, and SUFU; and Del/Dup: NF2 and SMARCB1	
NGS with reflex to Sanger as needed: BR. MAP2K1, MAP2K2, NRAS, PPP1CB, PTPN		Peripheral Nerve Sheath Tumor Testing		
SOS1, SOS2, and SPRED1; and Del/Dup: S	PRED1 and LZTR1	needed: NF1,	resh/Frozen Tumor for NGS with reflex to Sanger as NF2, KRAS, LZTR1, PTPN11 and SMARCB1; and Del/Dup: R1, and SMARCB1	
Sanger as needed: BRAF, CBL, HRAS, KRA			Tuberous Sclerosis Complex	
NF1, NRAS, PPP1CB, PTPN11, RAF1, RASA and SPRED1; and Del/Dup: NF1, SPRED1,		☐ TSC-NG: Fresh/Frozen Tumor or Tumor Block for NGS with reflex to Sanger as needed and Del/Dup: <i>TSC1</i> and <i>TSC2</i>		
	Additional I	nformation		
<u>Test Description Key:</u> Next Generation Sequencing (NGS) Sanger Sequencing (Sanger)	questions when completing this fo	rm.	via email at medgenomics@uabmc.edu if you have any w.uab.edu/medicine/genetics/medical-genomics-laboratory	

Deletion/Duplication analysis (Del/Dup)

Important points of consideration for testing

- The MGL offers next generation sequencing testing options that provide the ability to identify variants (indels and substitutions) as low as 3% of the alleles, depending on coverage in the regions of interest.
- NF1/SPRED1 biopsy-based testing is considered the "gold standard" approach for confirming a diagnosis of mosaic/segmental NF1 or Legius Syndrome.
- A minimum of two biopsies is required for NF1 testing. Two or more tumors are suggested for our other testing options. There are no additional fees associated with testing on additional biopsy specimens.
- When proceeding with tumor-based testing for NF2-related SWN, test code "SCH-NG" (NF2, SMARCB1, and LZTR1) is suggested unless the patient has findings unique to NF2.

Specimen Requirements					
Accepted Specimens	Specimen Information:				
Specimen requirements vary based on test requested; please see our website for more details.	□ Frozen □ Fresh				
more details.	□ Paraffin Curls □ Paraffin Block				
-CALs or Neurofibromas: require special media transport (kits are provided upon request, to be arranged <i>at least</i> one week in advance of procedure) -Fresh/Frozen Tumors: please submit a pathology report; for additional requirements, see tumor submission checklist -Formalin-Fixed Paraffin-Embedded Tumors (Tumor Block): please submit a pathology report; blocks are preferred to curls, when available; for additional requirements, see tumor submission checklist	□ Extracted DNA; Source:				
	☐ Biopsy-CAL-spot; # biopsies:				
	□ Biopsy-Neurofibroma; # biopsies:				
	Please note: failure to provide a date of collection can delay release of results				
	Tumor Collection Date (required):				

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Name: (First) (MI)	(Last)		DOB: (MM/DD/YY)			
Sanger Testing from Blood/Saliva/DNA						
NF1/Legius syndrome and Other RASo	pathy Related Conditions	Medium Chain Acyl-CoA Dehydrogenase Deficiency (MCADD)				
☐ NF1-R: Sanger and Del/Dup: <i>NF1 (RNA)</i>		☐ MCD1: Targeted analysis of exon 11 and, if needed, reflexive full gene				
☐ NFSP-R: Sanger and Del/Dup: NF1 (RNA)	and SPRED1 (gDNA)		y Sanger: ACADM			
Von Hippel-Lind	au	А	Autosomal Recessive Polycystic Kidney Disease			
☐ VHL1: Sanger and Del/Dup: VHL		☐ PKDL: Linka	age Analysis for informativity			
PTEN-Related Diso	rders					
☐ PTEN1: Sanger and Del/Dup: PTEN		☐ PKDPL: Pre	enatal Linkage (see Prenatal Specimen Requirements)			
Fragile X syndror	ne	FATHER: Na	ame and DOB (mm/dd/yyyy)			
☐ FRX: PCR and, if needed, reflexive confirm Southern blot analysis: <i>FMR1</i>	matory testing by	MOTHER: <u>N</u>	lame and DOB (mm/dd/yyyy)			
	Known Var	riant Testing				
and/or FISH analysis (Complete Previous Te	eviously identified known varia esting History: Page 1)	ant in any gene t	that is available at our lab by Sanger sequence, MLPA,			
	☐ KT2-NG: Targeted testing for a known variant with deep coverage of the alleles and detection of mosaicism for a variant present in at least 3% of alleles (Complete Previous Testing History: Page 1)					
☐ RT2: Targeted RNA-based testing for VO	US found during Next Generat	ion Sequencing	(Complete Previous Testing History: Page 1)			
□ PT2: Prenatal testing (see Prenatal Specimen Requirements; Complete Previous Testing History: Page 1) □ MCC: Blood specimen for mother provided for maternal cell contamination studies (required if not previously tested)						
☐ Other (unlisted options, please indicate below)						
Please contact lab before selecting this option						
		_				
	Additional	Information				
Test Description Key: Next Generation Sequencing (NG) Sanger Sequencing (Sanger) Deletion/Duplication analysis (Del/Dup)	Please contact the lab via phone questions when completing this	e (205) 934-5562 (s form.	or via email at medgenomics@uabmc.edu if you have any			
	Specimen R	equirements	s			
Accepted Prenatal Sp	ecimens		Prenatal Specimen Information:			
Specimen requirements vary based on test requested; please see our website		□ Amniotic fluic	d 🗆 Direct CVS (cleaned)			
for more detailsDirect CVS: minimum 10 mg cleaned villi		□ Cultured amn	niocytes Cultured villus cells			
-Direct amniotic fluid: minimum 10 ml fluid		Location of back-up culture (required):				
-Cultured CVS: Two T25 flasks (>70% confluent)		Sample Collected Date (required):				
-Cultured amniocytes: Two T25 flasks (>70% co	·	Specimen Information:				
Specimen requirements vary based on test requi		□ Peripheral Blo	lood (EDTA); # Tubes:			
for more details.		□ Extracted DN	, , ,			
-Blood: 3-6ml EDTA (must arrive within 60-72 hc RNA-based tests)	ours of collection for	□ Other, please describe:				
-DNA: extracted from lymphocyte cells, a minim	um of 25ul at 3μg, O.D. value at		ilure to provide a date of collection can delay release of results			
260:280nm ≥1.6 (must be extracted in a CLIA or -Sperm (for KT2-NG only): Fresh, sterile semental bank/cryobank facility			ected Date (required):			

LIFEMEDICAL GENOMICS LABORATORY

720 Twentieth Street South, Suite 330 Birmingham, Alabama 35294-0005

mingham, Alabama 35294-0005 Fax: (205) 996-2929 www.uab.edu/medicine/genetics/medical-genomics-laboratory

Phone: (205) 934-5562

UAB MGL Accession

Name:	(First)	(⋈1)	(Last)		DOB: (MM	/DD/YY)
			Bil	ling		
		☐ Please h	nold sample until furthe	er notice from the	ordering faci	lity.
			Important	Information		
	, ,	. , ,	agree that you have d		0.	
						e: If you are paying via self-payment or Requests for cancellation, test change, or
	ethod change of ongo	ing testing must	be submitted to the la	aboratory within t	hree working	days of specimen arrival. Individuals or
						rges for the cost of testing. nedical-genomics-laboratory
☐ Institution		trie billing polici	es is available at www	.uab.euu/meuicm	ie/genetics/n	iedical-genomics-laboratory
	Jilai bili	☐ Please cl	neck box if billing instit	ution should receiv	ve report dire	ectly
Institution:		- Fredde G	Teek box II billing Illotte	ation should recei	PO# (if applic	
IIISULUUOII.					РО# (п аррпс	abie).
Address:						
City:				State:		Zip:
Contact (Name and Title):						Preferred method of contact: ☐ Email ☐ Phone
Email:			Phone:			Fax:
☐ Self-Payr	ment Enclosed *PLE	ASE ENSURE ALL IN	IFORMATION IS LEGIBLE*	•		
	[☐ Visa	☐ MasterCard	☐ Discover	☐ Americ	can Express
Name as it app	pears on card:					
Card Number:				Expiration: (MM,	/YY)	3-digit Security Code:
Cardholder's Signature: Preferred method of contact: Email Phone						
Email:						Phone:
	Party Insurance Co					
Insurance Carr	ier:			_		rized in advance of shipment.
Insurance pre-verification/authorization previously performed? 🗆 Yes 🗖 No If yes, approval number is required:						
Please check box if you would <u>not</u> like insurance pre-verification/authorization to be performed by the MGL.						
ICD-10 Code	s (required):	Please send	a legible copy of the pa	tient's insurance ca	rd, front and b	ack.
Important Considerations for Insurance Billing						

For a list of contracted insurance companies, please visit our website or call our billing coordinator at 205-934-5523. As insurance prices are not listed, please call the billing coordinator to request a quote, if needed.

The MGL will contact the insurance provider to inquire regarding the CPT code coverage for all samples submitted for insurance payment. The healthcare provider will be contacted with the copay/deductible and also in cases where the insurance provider denies coverage of the requested codes or supporting documents are required from the provider to confirm coverage. The ordering provider/clinician's office is responsible for obtaining prior authorization, if it is required. This service is not offered for prenatal samples.

Please note: An insurance verification is not a guarantee of payment. Out of State Medicaid is not accepted under <u>any</u> circumstances. All RUSH fees must be paid up front. By completing this form, you agree that you have discussed the MGL's billing policies with your patient.

720 South Twentieth Street, Suite 330 Birmingham, Alabama 35294-0005 www.genetics.uab.edu/medgenomics

Tel: (205) 934-5562 Fax: (205) 996-2929

Informed Consent for Genetic Testing

This form does not need to be returned to the Medical Genomics Laboratory if Informed Consent portion of the Test Request form has been signed.*

I hereby consent for:						
Name:	DOB:	Gender:				

To participate in genetic testing for the following RNA/DNA-based cascade of tests ordered by my physician at the University of Alabama at Birmingham (UAB) Medical Genomics Laboratory (MGL):

Genetic Tests:	

I understand that:

- 1. Any biological samples submitted for genetic testing (e.g., blood, cheek cells, saliva, amniotic fluid, chorionic villi, tumor, and/or tissue) will be removed from me and/or my minor child(ren) using standard techniques which carry their associated risks.
- 2. Any samples obtained will be used for the purpose of attempting to determine if I and/or members of my family carry genetic changes in the disease genes ordered by my physician.
- 3. The genetic tests performed at the MGL are the most sensitive developed and are highly specific. However, sensitivity and specificity are test-dependent. Additional testing details and the specific detection rates of each test can be found at www.genetics.uab.edu/medgenomics.
- 4. The following are possible outcomes for the specific tests listed above:

Positive	Unknown Significance	Negative	
This is an indication that I may be predisposed to or have the specific disease, or condition tested. Further testing may be needed to confirm the diagnosis.	There may be a possibility that the laboratory findings will be ambiguous or of unknown significance. This may require additional testing from me or my family members. In rare circumstances, findings may be suggestive of a condition different than the diagnosis that was originally made.	technology and incomplete knowledge of genes, some changes in RNA/DNA or protein	

- 5. In other cases, the RNA/DNA test is unable to identify an abnormality although the abnormality may still exist. This event may be due to incomplete knowledge of the gene structure or an inability of current technology to identify certain types of mutations in the gene. When clinically necessary, the MGL may use a method called linkage analysis. This method is not a direct test, but will report the probability that you and/or family members have an inherited disease or disorder. In some families, the markers used in linkage analysis may be uninformative. If so, linkage testing cannot provide results for the family members in question.
- 6. The RNA/DNA analysis performed by the MGL is specific for the genetic test listed above and in no way guarantees my health or the health of my living or unborn children. The MGL cannot be responsible for an erroneous clinic diagnosis made elsewhere.
- 7. The tests performed at the MGL are expanded and improved continuously. The tests offered are not considered research but are considered the best and newest laboratory service that can be offered. Genetic testing is complex and utilizes specialized materials so there is always some very small possibility that the test will not work properly or that an error will occur. There is a low error rate (perhaps 1 in 1000 samples) even in the best laboratories. Additionally, in very rare instances, this test may reveal an important genetic change that is not directly related to the clinical reason for ordering this test. This would be considered an incidental finding. The MGL reserves the right to report these incidental findings if they are clinically relevant to the patients and/or their families In such instances, these results will be discussed with my healthcare provider and additional testing may be recommended. My signature acknowledges my voluntary participation in this test, but in no way releases the laboratory and staff from the MGL from their professional or ethical responsibility to me.
- 8. The MGL does not return DNA samples to individuals or physicians. While the MGL is not a specimen banking facility, in some cases it may be possible for the laboratory to reanalyze my remaining DNA upon request. The request for additional studies must be ordered by my referring physician/counselor and there will be an additional fee.

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- 9. An aliquot of my DNA/RNA may be used for validation, educational and/or research purposes. For some molecular genetic tests, a synopsis of clinical information and test results may be included in HIPAA-compliant, de-identified public databases as part of the National Institute of Health's effort to improve diagnostic testing and our understanding of the relationships between genetic changes and clinical symptoms. If I would like to opt out of participation, I can contact the MGL via email at medgenomics@uab.edu or calling the laboratory at 205-934-5562.
- 10. Because of the complexity of genetic testing and the important implications of the test results, results will only be reported to me through a physician, genetic counselor, or a certified genetics professional. The results are confidential to the extent allowed by law. They will only be released to other medical professionals or other parties with my written consent or as otherwise allowed by law. Participation in testing is completely voluntary.
- 11. For Prenatal Testing: If prenatal diagnosis is being performed, fetal cells obtained by chorionic villus sampling or amniocentesis will be used. In order to perform accurate prenatal diagnosis, biological samples are required for the fetus as well as from the affected individual in the family and from the biological mother.
- 12. I and/or my physician/counselor have signed the informed consent portion of the test order form indicating that we have discussed the items on this document and I will also receive a copy of this consent form.

Subject's Signature	Date	Physician's Signature	Date
Please Print Subject's Name		Please Print Physician's Name	
Assent of Parent	Date	Genetic Counselor's Signature	Date
Assent of Child	 Date	Please Print Genetic Counselor's I	Name



MEDICAL GENOMICS LABORATORY: TSC1/TSC2 PHENOTYPIC CHECKLIST FORM

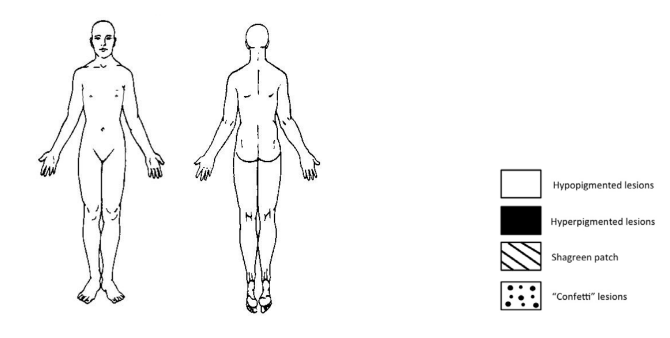
Patient Name:	□Male □Female	Date of Birth / /
Referring Physician:		Date of Exam//
Thank you for completing this form; phenotypic information may im	prove our ability to in	terpret your results.
DEMOGRAPHIC INFORMATION Ethnicity: ☐ White ☐ Black ☐ Native American ☐ Hispanic ☐ Asian	Other:	
DIAGNOSIS		
2012 International TSC Consensus Conference Clinical Criteria: (r (Criteria are listed below with major criteria marked by # and n Definite TSC (2 major or 1 major plus 2 minor feature) Possible TSC (1 major or 2 or more minor features) Does not meet TSC Criteria	ninor criteria marked	
Clinical Concern for Mosaicism: ☐ No ☐ Yes Family history: ☐ Sporadic ☐ Familial ☐ Unknown		
Known Familial Mutation: No Yes (provide information:)
Familial cases: Please provide pedigree and details on the affection status of analysis if available.	family members on a sep	parate page. Attach prior mutational
SIGNS AND SYMPTOMS (Major criteria marked by # and mi	nor criteria marked	by *.)
2) "Confetti" skin lesions: 3) Facial Angiofibromas: 4) Shagreen Patch: 5) Cephalic Fibrotic Plaque: 6) Ungal/Periungal fibromas: None Present* None Present* None Present* None Present* None Present* None Present* 1-3 >3# [None Present* None Present* 1-2 >2# [None	Unknown Unknown Unknown Unknown Unknown Unknown Unknown Unknown	
A digital picture of the skin findings would be very helpful.	manigo en page el	
Neurological (Imaging) 8) Cortical Dysplasia: Cortical Tubers: Cerebral White Matter Radial Migration Lines: None 9) Subependymal nodule (SEN): None 10) Subependymal Giant Cell Astrocytoma (SEGA): Histopathologically Confirmed?	☐ Present# ☐ Present# ☐ Present# ☐ Present# ☐ Present# ☐ Yes	☐ Unknown ☐ Unknown ☐ Unknown ☐ Unknown
Neurological/Psychiatric (Clinical) 11) Seizures: ☐ None ☐ Present (Describe type, if known:	☐ Present ☐ Unknow ☐ ADHD ☐ Isolated ☐ unknown ☐ Other	hyperactivity Aggression
Renal 14) Angiomyolipomas: Histopathologically Confirmed? Is a malignant angiomyolipoma present? No 15) Renal Cell Carcinoma: None 16) Renal Epithelial Cysts: None 17) Polycystic Kidney Disease Features: None	☐ 1-2	Unknown Unknown Unknown

MEDICAL GENOMICS LABORATORY: TSC1/TSC2 PHENOTYPIC CHECKLIST FORM



Patient Name:				Date of B	sirth//
(Criteria are listed below w	ith major criteri	a marked by	# and minor	criteria marke	d by *.)
Pulmonary 18) <u>Lymphangioleiomyomatosis</u> Histopathologically Confirm		☐ None ☐ No	☐ Present#	Unknown	
Cardiac 19) <u>Rhabdomyomas:</u>	☐ None ☐ Prese	nt [#]	y/Prenatally pres	sent, but regressed	Unknown
Dental 20) <u>Dental Enamel Pits:</u> 21) <u>Intraoral Fibromas:</u>		☐ 1-3 ☐ >3* ☐ 1-2 ☐ >2*	Unknown (Location:)
Ophthalmological 22) Retinal Hamartomas: 23) Retinal Achromic Patch: 24) Retinal Astrocytic Hamarton	☐ None ☐ None ☐ None	☐ Sing ☐ Pres ☐ Pres	ent* 🔲 Un	ıltiple [#] □ Unk ıknown ıknown	known
Neuroendocrine 25) <u>Neuroendocrine Tumors</u> :	☐ None ☐ Unkn	own 🗌 Yes (S	pecify type:)
Other 26) Nonrenal Hamartomas: 27) Additional Phenotypic Inform	☐ None ☐ Presenation:	ent* 🗌 Unkno	wn		

Indicate location/size of hypomelanotic macules or other dermatological lesions \downarrow



www.genetics.uab.edu/medgenomics

Tumor Specimen Submission Checklist

The following requirements must be met in order to process tumor specimens. The UAB Medical Genomics Lab now proudly offers Fresh/Frozen/FFPE Tumor testing utilizing Next Generation Sequencing. Please confirm that the submitted tissue meets each requirement by placing a check mark next to each statement. If your pathology department is unable to confirm this information for you, please select the check-box below. If Neurofibromatosis Type 1 is your primary concern, neurofibromas and biopsied CALs still require being collected in our media and are run via Sanger sequencing. Please contact the MGL at (205) 934-5562 to request collection media for these sample types.

For NGS Sequencing

	1 0		
	Fresh Tumor Specimen Checklist		
	This tumor is at least 5mm-cubed		
	This specimen contains at least 60% pure tumor content		
Each specimen has been sent in individual vials of basic, sterile culture media such as RPMI or PBS			
	and is marked with its specific location and/or tumor type		
	Frozen Tumor Specimen Checklist		
	This tumor is at least 5mm-cubed		
	This specimen contains at least 60% pure tumor content		
	This specimen has been snap frozen and sent on dry ice and is marked with its specific location		
	and/or tumor type		

	Formalin-fixed paraffin embedded block			
	This tumor block has a surface area of a least 5mm squared <u>or</u> This specimen contains at least 3-6			
	loose paraffin curls (no slides) that are 30-50 microns thick			
This tumor specimen contains greater than 70% nucleated cells				
	This specimen contains at least 60% pure tumor content			
	Notes or Special Comments			