



Interpace Diagnostics 2515 Liberty Avenue Pittsburgh, PA 15222

MOLECULAR REQUISITION FORM

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Phone: 844-405-9655 Email: labsupport@ir	iterpace.com		interpace.com
(1) Patient Information Please print or adhere patient label. Must include two (2) unique identifiers.	Chani	2 Physician	
		tting Physician	Referring/Treating Physician
Last Name:First Name:	Account #: Office/Hospital:		Account #: Office/Hospital:
Date of Birth (mm/dd/yy):/	Address:		Address:
	Phone:		Phone:
SSN/MRN: Gender: \square M \square F	Fax: Office Contact:		Fax: Office Contact:
(3) Billing Information	Email:		Email:
Procedure Location:			Contact Preference:
☐ Outpatient ☐ Non-Hospital Affiliated Setting ☐ Private Practice ☐ Inpatient/Discharge Date: / /	Institution NPI:		☐ No Contact ☐ CC Test results
ICD CODE:	Physician NPI:		
Codes for your consideration (please do not circle, see reverse side for more information):			Institution NPI:
E04.2 Nontoxic, multinodular goiter thyroid gland E04.1 Nontoxic, single thyroid nodule			Physician NPI:
D44.0 Neoplasm of uncertain behavior of thyroid gland D34.0 Benign neoplasm of thyroid			
The diagnosis code(s) provided should always be based upon what can be supported within the patient's medical records. Testing cannot be done unless ICD code(s) are included.			
(4) Specimen & Diagnosis Information			
Please indicate type and number submitted			
Submitted Specimen(s):			
# FNA in Collection Buffer Vial(s)		(5) Method of Pay	ment Information
# Cytology Slide(s)	A COPY OF THE	PATIENT'S BILLING AND DEN	OGRAPHICS INFORMATION IS REQUIRED
Submitted Collection Date: / /			INFORMATION WILL DELAY RESULTS.
For multiple nodules, indicate the locations on the diagram and correlate with labels	☐ Medicare ☐ M	1edicaid □ Private Insurance	e □ Ordering Institution □ Self Pay
attached below.	Interpace Diagnos	tics will bill directly for insured p	atients, wherever permitted by government
A B		= -	rangements. If patient or insurance information
Size: Size:	is not completed t	or attached, your facility will be b	mieu.
A A B A			
	(6) Test Menu and Authorization		
$(\circ \circ \circ) \qquad (\circ \circ \circ)$		<u> </u>	
	☐ ThyGeNEX	T® w/ Reflex to ThyraM	IR®v2
Right Left Right Left	, ,	,	iscriminates benign from malignant nodules
Cytology Diagnosis (Bethesda Category):	•		V600E and other mutations (BRAF-like
A B A B			nancy are included in ThyGeNEXT. Also cer (<i>RAS</i> -like mutations) are also included in
☐ ☐ Atypical/FLUS (III) ☐ ☐ Nondiagnostic (I) ⁺	ThyGeNEXT. If m	nutations in ThyGeNEXT are n	egative or not fully predictive of
□ □ Suspicious for Neoplasm (IV) □ □ Benign (II)+	malignancy, Thyr	aMIRv2 testing will be perfor	med in reflex.
□ Suspicious for Cancer (V) □ □ Malignant (VI) ⁺ *Requires Letter of Medical Necessity (LOMN)	☐ ThyGeNEXT	Γonly	
PLEASE ATTACH A COPY OF THE CYTOLOGY REPORT			
		•	or which reimbursement from Medicare or third-
Ultrasound Characteristics (check all that apply):		_	ally necessary for the diagnosis, care, and providing this patient's test results to the
A B	patient's third-party	payor. I certify that the treating	physician has ordered the above test.
□ □ Peripheral Vascularity □ □ Rim Calcifications	MD/DO Signature	2	
☐ ☐ Intranodular Vascularity ☐ ☐ Macrocalcifications		- 	_
□ □ Avascular □ □ Microcalcifications	Print Name:		
☐ ☐ Hyperechoic ☐ ☐ Isoechoic	Order Date:		
☐ ☐ Hypoechoic			
Clinical History/Comments:	<u> </u>		
Specimen A Specimen A	Specimen	В	Specimen B
Button Many	D-11		Dationt Name
Patient Name: Patient Name: DOB: DOB:	Patient Nam DOB:	e:	Patient Name: DOB:
, : 000,	,		



Incomplete or incorrect information will lead to testing delays

Section 1. Patient Information

All fields are required. Include patient first and last name, date of birth, SSN/MRN code, and sex. A complete patient label can be adhered in this field as long as patient information is complete and includes 2 unique identifiers. Incomplete information will result in testing delays.

Section 2. Physician Information

This section may be pre-filled if an account has been set up with Interpace Diagnostics. If blank, complete each field. Account Number should match your Interpace Diagnostics account number (leave blank if unknown). Complete physician institution contact information. Please avoid Office/Hospital acronyms or abbreviations and spell out the institution name.

Referring/Treating Physician: Please provide full name and phone/fax number for referring/treating physician, if applicable. Check the box to indicate if you would like the referring/treating physician to receive a copy of the results. Leaving contact information incomplete will result in processing delays.

Section 3. Billing Information

Check the box to indicate where procedure was performed (Non-hospital affiliated setting, Private Practice, Outpatient, or Inpatient with discharge date).

Testing cannot be performed unless ICD-10 code(s) are included.

Write in the appropriate ICD-10 code based on the patient's medical records. The ICD diagnosis code must be defined by the most detailed level of specificity available. If a diagnosis code cannot be supported by the patient's medical record, then the code should not be used for ordering laboratory services. The list of common ICD diagnosis codes for the characterization of thyroid nodules shown below is not complete. Please refer to the ICD manual for a complete listing.

Common ICD-10 Diagnosis Codes:

D34.0- Benign neoplasm of thyroid gland

D44.0-Neoplasm of uncertain behavior of thyroid gland

D44.9-Neoplasm of uncertain behavior of unspecified endocrine gland

E01.0-lodine-deficiency related diffuse (endemic) goiter

E01.1 Iodine-deficiency related multinodular (endemic) goiter

E01.2-Iodine-deficiency related (endemic) goiter, unspecified

E04.0-Nontoxic diffuse goiter

E04.1-Nontoxic single thyroid nodule

E04.2-Nontoxic multinodular goiter

E04.8-Other, specified nontoxic goiter

E04.9-Nontoxic goiter, unspecified

Section 4. Specimen & Diagnosis Information

Submitted Specimen(s): Check the box to indicate type of specimen being sent in for testing OR indicate number of slides being sent (REQUIRED). If submitting multiple nodules, indicate type and quantity separately.

Specimen Collection Date: Enter the date of the procedure when specimen was collected. Mark the circle on the thyroid diagram to indicate location of the nodule. The requisition can be used to submit specimens for up to 2 locations. Use letter A for the first specimen and letter B for the second. Provide a descriptive name for each location on the line provided along with the size of each nodule.

Cytology Diagnosis (Bethesda Category): Check the box that corresponds with the patients' medical records to indicate the cytology diagnosis. A copy of the corresponding cytology report is requested to be sent with this specimen, as available. If these reports are not available at time of specimen submission, please forward to Client Services (fax: 1-888-674-6894 or 412-224-6425) when received.

Section 4. Specimen & Diagnosis Information (cont.)

Please note that for Bethesda I, II, and VI categories a Letter of Medical Necessity is required to start testing.

Ultrasound Characteristics: Indicate key characteristics identified from patient's medical records & ultrasound report. Provide relevant clinical history for the patient. Please provide relevant medical records at no cost when requested by the patient's insurance carrier for reimbursement.

Section 5. Method of Payment Information

Check the box indicating how the testing should be billed. Interpace Diagnostics will submit claims to all private insurance, Medicare, and other government plans for insured patients.

Patient Contact Information: Please provide a copy of the patient's face sheet or demographics page to include the patient's full name, gender, date of birth, address, and phone number.

A copy of the patient's billing information MUST be submitted with specimen.

Medicare/Medicaid/Private Insurance: Provide a clear copy of the front and back of the patient's primary insurance/Medicaid/other payer card. If the patient has a secondary insurance please provide a clear copy of the front and back of the secondary insurance card.

Ordering Institution: Check this box if Interpace Diagnostics is to bill the ordering institution for the ordered testing.

Patient Self-Pay (no insurance): Check this box if the patient has no insurance.

Section 6. Test Menu and Authorization

Specimen processing cannot begin until there is a clear indication of the type of testing to be performed (check box). Please indicate tests requested for patient specimen. Details of each test can be found on our website and in test details from our Directory of Laboratory Services. Only 1 test should be selected.

The ordering physician is required to sign and date the requisition as authorization for Interpace Diagnostics to perform testing. Stamped signatures or physician initials cannot be accepted. An incomplete signature will result in testing delays. Order date cannot precede Collection Date of the specimen. **Post-dated requisitions are not accepted.**

Questions?

Contact Interpace Diagnostics Customer Service at:

Phone: 844-405-9655 or 412-224-6900 Fax: 1-888-674-6894 or 412-224-6425 Email: clientservices@interpace.com

Website: www.interpace.com

