

Patient Information					
Last Name:					
First Name:	MI: Sex:□M	□F			
DOB (mm/dd/yyyy):	SSN:				
Parent/Guardian (if applicable):(REQUIRED if patient is <18 years old at time of sample collection)					
Address:	.ddress:City:				
State: Zip: Phone:					
ICD-10 Diagnosis Code(s)					
Primary ICD-10 Code (REQUIRED)	Additional ICD-10 Code(s) (OPTIONAL)				
1	2 3 4				
	Billing Information				
BILL: Insurance*	Laboratory Patient/Self-Pay Provider Acco	unt			
*Please provide details be	low <u>AND</u> attach a front/back copy of insurance care	d(s).			
Insurance Carrier:	Policy #:				
Pro	ovider/Account Information				
Account Name/Address					
Phone:	Fax:				
Ordering Provider Selection & Signature (REQUIRED)					
	ordering provider from list below (circle or √). ted, please provide their FULL NAME AND NPI:				
If provider is not lis	ted, please provide their FULL NAME <u>AND</u> NPI:				
If provider is not lis Provider: I certify the ordered test(s)	ted, please provide their FULL NAME AND NPI:	osis,			
Provider: Certify the ordered test(s) care, and/or treatment of ti	NPI: is/are reasonable and medically necessary for the diagnosis patient's condition as documented in the medical reco	ord.			
Provider: I certify the ordered test(s) care, and/or treatment of t if this patient is <18 years informed that they will be	NPI: is/are reasonable and medically necessary for the diagnosis patient's condition as documented in the medical recoold at the time of testing, the parent/guardian has be responsible for any costs associated with testing.	ord. een			
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IBD: Inflammatory bowel disease; UC: Ulcerative colitis; CD: Crohn's disease; IFX: Infliximab; ATI: Antibodies-to-IFX; ADA: Adalimumab; ATA: Antibodies-to-ADA

☐ No Results to Lab

Results:

☐ Mail ☐ Fax

All information requested is REQUIRED to process testing and/or submit insurance claims on your patient's behalf.

Prometheus Client Services | 888-423-5227 | 6:00 am - 4:30 pm Pacific Time

Select Test(s) To Be Performed									
— Diagnosis & Risk Stratification —									
■ IBD sgi Diagnostic®† (#1800) Aids in diagnosis and differentiation of IBD from non-IBD and UC from CD. — Conditional testing if IBD sgi result is indicative of IBD —									
Pat NO	D Respondr® T tient Weight in TE: Patient mus — Conditional	kg (REQU t be ≥35 k	JIRED*) g at tim	e of sa	mple c	olle	ction.		2)
NO	D Monitr® Croh TE: Patient mus D Crohn's Prog	t be ≥12 ye	ears old	,	e of sar	mple	e colled	ction.	
☐ Mon	nitr® Crohn	's Disea	ase (#	,		pat	tients ≥	≥12 vears o	old.
Measure and track endoscopic disease activity in CD patients ≥12 years old. ☐ Crohn's Prognostic [†] (#2001) Evaluate probability of disease progression in Crohn's disease patients.									
	_	· Therap	eutic S	Selecti	ion —				
— Therapeutic Selection — ☐ Respondr® TNF† (#4200) Indicates likelihood of anti-TNF response, prior to initiation of therapy, by assessing pre-therapy drug clearance and genetic predisposition to immunogenicity.									
	Patient Wei ight in kg = lbs ÷ 2.2)								
	_	Precision	n-Guid	ed Do	sing ·				
PredictrPK® IFX Induction (#3500) Predicts IFX levels at 4-, 6- and 8-weeks post induction dose 3; calculates IFX clearance and measures serum IFX, ATI and albumin in IBD patients. Sample collection: ≤3 days prior to IFX induction dose 3. IFX Induction Dosing (REQUIRED)*									
	Infusion [(mm/dd/s	ate	Tota	al Dose .g. 485	in mg			eight in kg = lbs ÷ 2.2	
Dose 1					m	ng			kg
Dose 2					m	ng			kg
Dose 3*					n	ng			kg
*Scheduled	l infusion date, es	imated tota	al dose a	nd antio	cipated	pati	ient wei	ight.	
PredictrPK® IFX Maintenance (#3400) Predicts IFX trough levels with current & alternative dosing; calculates IFX clearance and measures serum IFX, ATI and albumin in IBD patients. Sample collection: ≤3 days prior to week 14 or, after ≥14 weeks of IFX therapy, anytime ≥20 days after any maintenance infusion, up to and including at trough. Last Administered IFX Infusion (REQUIRED)*									
Interval (we]5 []6	□ 7	□ 8	<u>□</u> 9			Other:	
	Last Infusion n/dd/yyyy)		al Dose e.g. 485					ght in kg lbs ÷ 2.2)	
					mg				kg
PredictrPK® ADA Maintenance (#3700) Predicts ADA levels with alternative dosing; calculates ADA clearance and measures serum ADA, ATA and albumin in IBD patients ≥12 years old. Sample collection: Anytime within the injection interval after ≥8 weeks of ADA therapy. Current ADA Dosing (REQUIRED)*									
	Last Injection	Tent ADA	Dose		DIKED	<u>)</u>		terval	
(mm	n/dd/yyyy)	Г	(mg) (days) □40 □80 □7 □14 □Other:						
Miscellaneous Testing									
Other Prometheus Testing†: †Acknowledgment of informed consent required if genetic testing is indicated above. Refer to page two or contact Client Services to confirm specimen requirements.									
	— Snasim	en collecti	on roau	iromon	tc on n	200	2 _		

[†]Acknowledgment of informed consent for genetic testing required. ^{*}Failure to provide the required data, at the time of sample collection, may result in reporting delay/cancellation.

Specimen Collection and Handling Procedures						
Test Ordered	Turnaround Time ^a	Transport Kit and Storage Conditions	Specimen Type and Volume	Specimen Collection Tube	Specimen Stability	
7C4 Diagnostic	7 days	Cold pack REQUIRED Do not freeze	1.0 mL Serum	Serum Separator Tube or Red-Top Tube	Room temp: 3 days Refrigerated: 7 days	
Anser* ADA, IFX, RZB, UST, VDZ	3 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum (0.5 mL for Peds)	Serum Separator Tube or Red-Top Tube	Room temp: 14 days Refrigerated: 14 days	
Celiac PLUS [†]	4 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum AND 2.0 mL Whole Blood	Serum Separator Tube or Red-Top Tube AND EDTA/Lavender-Top Tube	Room temp: 7 days Refrigerated: 30 days	
Celiac Genetics†	4 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Whole Blood	EDTA/Lavender-Top Tube	Room temp: 7 days Refrigerated: 30 days	
Celiac Serology	4 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum (0.5 mL for peds)	Serum Separator Tube or Red-Top Tube	Room temp: 7 days Refrigerated: 30 days	
Crohn's Prognostic†	7 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum AND 2.0 mL Whole Blood	Serum Separator Tube or Red-Top Tube AND EDTA/Lavender-Top Tube	Room temp: 7 days Refrigerated: 7 days	
IBD sgi Diagnostic†	4 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum AND 2.0 mL Whole Blood	Serum Separator Tube or Red-Top Tube AND EDTA/Lavender-Top Tube	Room temp: 7 days Refrigerated: 21 days	
Monitr Crohn's Disease	5-7 days	Cold pack preferred <u>Do not freeze</u>	2.0 mL SPUN Serum	SPUN Serum Separator Tube	Room temp: 3 days Refrigerated: 14 days	
PredictrPK IFX, ADA	3 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Serum (0.5 mL for Peds)	Serum Separator Tube or Red-Top Tube	Room temp: 14 days Refrigerated: 14 days	
Respondr TNF†	4 days	Ambient or cold pack <u>Do not freeze</u>	3.0 mL Whole Blood	EDTA/Lavender-Top Tube	Room temp: 10 days Refrigerated: 30 days	
Thiopurine Metabolites	3 days	Cold pack preferred <u>Do not freeze</u>	5.0 mL Whole Blood	EDTA/Lavender-Top Tube	Room temp: 24 hours Refrigerated: 8 days	
TPMT Enzyme	3 days	Cold pack preferred <u>Do not freeze</u>	5.0 mL Whole Blood	EDTA/Lavender-Top Tube	Room temp: 24 hours Refrigerated: 8 days	
TPMT Genetics†	4 days	Ambient or cold pack <u>Do not freeze</u>	2.0 mL Whole Blood	EDTA/Lavender-Top Tube	Room temp: 10 days Refrigerated: 30 days	

^aBusiness days from date of receipt.

Specimens should be labeled with 2 identifiers and date of collection. Examples of acceptable identifiers include, but are not limited to: patient name, date of birth, hospital number, requisition, accession, or unique random number. Unlabeled specimens will not be accepted for testing.

SHIPPING INSTRUCTIONS: Prometheus has an agreement with FedEx* for express overnight delivery within the United States and Canada. Please call FedEx at 1-800-GoFedEx (463-3339)

to schedule a pickup. FedEx will pick up your specimens and ship them to Prometheus Laboratories Inc. in San Diego at no expense to you. Prometheus will provide specimen transportation kits upon request. NOTE: Multiple specimens may be shipped in a single transportation kit. For more information, call Client Services at 888-423-5227, or go to prometheuslabs.com.

Important Ordering Information





Respondr and PredictrPK require specific patient and treatment inputs for processing and resulting. You may need to request this data from the infusion center.

Failure to provide the required data, at the time of ordering, may result in reporting delays and/or possible cancellation.

Weight in kg

- Patient's weight, in kilograms (kg), at the time of sample collection (Respondr) or at the designated infusion (PredictrPK IFX)
- Formula for reference: kg = lbs ÷ 2.2

Total Dose in mg

- Total amount of IFX administered, in mg (e.g. "485 mg") Note: We CANNOT accept derived dosing such as "5 mg/kg"
- If not all IFX in a vial is given, estimate the total dose given to the nearest 5 mg

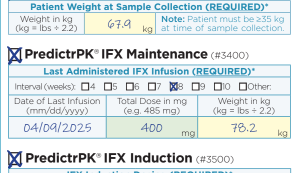
Dose 3 Data (PredictrPK IFX Induction)

As sample collection occurs prior to infusion/dose 3, please provide:

- · Scheduled date of infusion 3
- Anticipated total IFX dose, in mg, to be administered
- Estimated patient weight, in kg, at infusion 3

Example orders are provided for illustrative purposes only.

Respondr® TNF† (#4200)



A licalculate in X illadoction (#5500)						
IFX Induction Dosing (REQUIRED)*						
	Infusion Date (mm/dd/yyyy)	Total Dose in mg (e.g. 485 mg)	Weight in kg (kg = lbs ÷ 2.2)			
Dose 1	07/18/2025	250 _{mg}	42.7 kg			
Dose 2	08/01/2025	250 mg	43.1 _{kg}			
Dose 3*	08/29/2025	350 mg	44.2 kg			

REQUIRED ACKNOWLEDGMENT OF INFORMED CONSENT FOR GENETIC TESTING

I warrant that this test was ordered and that I have obtained the appropriate prior written consent. This written consent was signed by the person who is the subject of the test (or if that person lacks capacity to consent, signed by the person authorized to consent for that person) and includes the following (unless certain that the following information is not required by the state in which I practice): The purpose of this test is to determine if the patient may have a variant in the gene(s) being tested, which has been found to be associated with this condition.

- This test will only test for this specific condition and will not detect ALL possible variants within this gene, nor will it detect variants in other genes.
- Prior to my signature of the written consent, a qualified healthcare professional discussed with the patient the genetic test ordered and described the steps involved in the test, the constraints of the procedure, and its accuracy.
- 4. My patient was advised by a qualified healthcare professional of the risks and benefits of genetic testing, and advised of the significance of a positive and negative result.
- The patient understands that a positive test result is an indication that the patient may be predisposed to, or have, the condition listed.
- The patient understands that the test may fail, that the results may be non-informative or not predictive for his/her case, and that the test may reveal information that is unrelated to their intended purpose
- No unauthorized test is performed on specimens.
- NY-state specimens will be destroyed within 60 days after collection when no longer required for clinical purposes.
- The informed consent does not authorize the use or release of any other identified medical information unrelated to this genetic test.
- 10. The patient understood that he/she could see professional genetic counseling prior to undergoing the testing procedure, and received written information identifying a genetic counselor or medical geneticist by his/her treating provider.

Prometheus tests are laboratory-developed tests that were developed, and analytically and clinically validated by Prometheus Laboratories Inc. under federal Clinical Laboratory Improvement Amendments (CLIA) guidelines, and are performed exclusively in our high complexity CLIA certified and College of American Pathologists accredited clinical laboratory. As laboratory developed tests, they have not been cleared or approved by the US FDA. The tests may be covered by one or more US pending or issued patents - see prometheuslabs.com/patents. Prometheus, Anser, IBD sgi Diagnostic, Monitr, PredictrPK and Respondr are registered trademarks of Prometheus Laboratories Inc, San Diego, California. All other trademarks or service marks are the property of their respective owners. ©2025 Prometheus Laboratories Inc. All rights reserved.

[†]Acknowledgment of informed consent for genetic testing required.