

# Helpful Hints for Ordering the clonoSEQ® Assay

Questions? Contact Adaptive Clinical Services  
P: 888 552 8988 | F: 866 623 4408  
clinicalservices@adaptivebiotech.com

clonoSEQ.com



## SPECIMEN PREPARATION

### Review Requirements

- ✓ Guidance for sending specimens can be found on [clonoSEQ.com](https://clonoSEQ.com).

### Provide Ample Material

- ✓ If ordering more than one Clonality Test (i.e. B-cell **and** T-cell\*) include double the sample material requested in the Specimen Requirements document.

### Label w/ 2 Unique Identifiers

- ✓ Each specimen must be labeled with **2 unique patient identifiers**. These identifiers must match the patient information you entered on the **Test Requisition Form (TRF)**.



## TEST REQUISITION FORM (TRF)

### Complete



#### ✓ FILL OUT SPECIMEN FIELDS

Fully complete the Specimen Information section of the TRF, especially the "collection date." These fields are not required to save and print a TRF, but **they are required prior to sending a specimen**.

### Verify



#### ✓ VERIFY INSURANCE INFORMATION HASN'T CHANGED

If you previously provided a patient's insurance details, they will automatically appear when you start a new TRF. Verify this information is correct before saving and printing the TRF.

### Sign



#### ✓ PROVIDE SIGNED & DATED TRF

Completed TRF must be sent to Adaptive before a sample can be processed.

### Request



#### ✓ PROVIDE COMPLETE PATHOLOGY REPORT (ID SPECIMENS ONLY)

If you would like assistance from Adaptive to request a pathology sample, **send a copy of the patient's complete pathology report with the signed and dated TRF** to Clinical Services.

#### ✓ PROVIDE COPY OF INSURANCE CARD

If you selected "Insurance" as the billing type on the TRF, include a front-and-back copy of the patient's insurance card with the specimen shipment.



Required fields are highlighted in yellow on a printed TRF as a reminder to complete them before sending. **TRFs with missing or incorrect information will delay ordering and the delivery of patient results.**

clonoSEQ®

Copyright © 2021 Adaptive Biotechnologies Corp.  
All rights reserved.  
PM-US-cSEQ-0112-3 | MRK-00195-4

\*T-cell testing is available via Adaptive's CLIA-validated LDT service and has not been cleared or approved by the FDA.

clonoSEQ® is available as an FDA-cleared *in vitro* diagnostic (IVD) test service provided by Adaptive Biotechnologies to detect minimal residual disease (MRD) in bone marrow from patients with multiple myeloma or B-cell acute lymphoblastic leukemia (B-ALL) and blood or bone marrow from patients with chronic lymphocytic leukemia (CLL). clonoSEQ is also available for use in other lymphoid cancers and specimen types as a CLIA-validated laboratory developed test (LDT). For important information about the FDA-cleared uses of clonoSEQ including test limitations, please visit [clonoSEQ.com/technical-summary](https://clonoSEQ.com/technical-summary).

Adaptive  
biotechnologies