

Instructions for Use

FLT3 ITD MRD Assay

For detection of internal tandem duplication (ITD) mutation in the FMS-like tyrosine kinase 3 (FLT3) gene.



RUO For Research Use Only. Not for use in diagnostic procedures.



Catalog # **Products** Quantity

Table of Contents

| 1. | Assay Use | | | | | |
|------------|--------------|------------------------------------------------------------------------------|----------|--|--|--|
| 2. | SUMMAR | Y AND EXPLANATION OF THE TEST | 3 | | | |
| | 2.1. | Summary | 3 | | | |
| | 2.2. | Explanation | 3 | | | |
| 3. | PRINCIPI | ES OF THE PROCEDURE | 3 | | | |
| J . | | | | | | |
| | 3.1. | PCR Amplification of the Internal Tandem Duplication (ITD) Mutations of FLT3 | | | | |
| | 3.2. | Next-Generation Sequencing (NGS) | | | | |
| | 3.3. | Sample Multiplexing | | | | |
| | 3.4. | Measurable Residual Disease Evaluation | | | | |
| 4. | REAGENT | S AND MATERIALS | 5 | | | |
| 5. | SOFTWAR | RE | е | | | |
| | 5.1. | FLT3 ITD MRD v1.2 Software | е | | | |
| 6. | WARNIN | GS AND PRECAUTIONS | 7 | | | |
| | 6.1. | Cyber Security Precautions | | | | |
| 7. | SPECIME | N COLLECTION AND PREPARATION | 8 | | | |
| | 7.1. | Precautions | 8 | | | |
| | 7.2. | PCR-Interfering Substances | | | | |
| | 7.3. | Specimen Requirements and Stability | | | | |
| | 7.4. | Sample Preparation | | | | |
| 8. | Assay Pr | ROCEDURE | <u>ç</u> | | | |
| | 8.1. | PCR Setup | c | | | |
| | 8.2. | First PCR Product Purification (0.8x Cleanup) | | | | |
| | 8.3. | Second PCR Product Purification (0.7x Cleanup) | | | | |
| | 8.4. | Quantification and Validity of Purified PCR Products | | | | |
| | 8.5. | Pooling and Quantification of the Library | | | | |
| | 8.6. | Preparation of the Library Pool for the MiSeq™ Sequencing Run | | | | |
| | 8.7. | Preparation of the MiSeq™ Reagent Cartridge | | | | |
| | 8.8. | MiSeq™ Sample Sheet Creation | | | | |
| | 8.9. | MiSeq™ Reagent Cartridge Loading | | | | |
| | 8.10. | MiSeq [™] Run Start | | | | |
| | 8.11. | Acceptance Criteria | | | | |
| 9. | DATA AN | ALYSIS | 16 | | | |
| | 9.1. | FLT3 ITD MRD v1.2 Software | 16 | | | |
| | 9.2. | Sample Interpretation | | | | |
| | 9.3. | ITD and Insertion Variant Detection | 16 | | | |
| | 9.4. | Sequence Annotations | 16 | | | |
| 10. | TROUBLE | SHOOTING | 17 | | | |
| 11. | REFEREN | CES | 17 | | | |
| 12. | TECHNICA | AL AND CUSTOMER SERVICE | 17 | | | |
| Con | TACT INFO | RMATION | 17 | | | |
| 13. | SYMBOLS | | 18 | | | |
| 14. | LEGAL NOTICE | | | | | |

1. Assay Use

This Research Use Only assay targets the juxtamembrane (JM) region of the FLT3 gene to identify ITD mutations.

2. Summary and Explanation of the Test

2.1. Summary

The FMS-like tyrosine kinase 3 (*FLT3*) gene encodes a receptor tyrosine kinase that is normally expressed on many cell types including hematologic stem cells. Mutation of the *FLT3* receptor, by internal tandem duplication (ITD) of the juxtamembrane domain, causes constitutive activation of the *FLT3* receptor. *FLT3* ITD mutations are present in about 25% of patients with Acute Myeloid Leukemia (AML) and are characterized by an increased risk of relapse. Since *FLT3* ITD mutations are present in 75% of relapsed subjects which were *FLT3*-ITD mutated at diagnosis², ITDs can serve as a biomarker for monitoring. The ability to sensitively and specifically detect measurable residual disease (MRD) in leukemia subjects has proven to be useful both in the clinical management of select subtypes of the disease.

2.2. Explanation

The *FLT3* ITD MRD assay is designed to detect the presence of ITD mutations within exons 14/15 of the *FLT3* gene in a given human DNA sample. The test is an amplicon-based approach which uses polymerase chain reaction (PCR) to amplify the region of interest and next-generation sequencing (NGS) to detect the region of interest. Primers included in the master mixes are designed with Illumina® adapters containing unique indexes. This method allows for pooling of amplicons from several different samples onto one MiSeq™ flow cell, allowing for up to 24 samples to be analyzed in parallel in a single run.

The assay includes twenty-four dual-indexed PCR master mixes, along with positive and negative controls. The *FLT3* ITD MRD assay was designed with 24 unique index combinations, labeled as IA or IB.

The associated RUO FLT3 ITD MRD v1.2 software provides a simple and streamlined method of analysis.

3. Principles of the Procedure

05 03 04 DNA PCR NORMALIZING. SEQUENCING FIT3 ITD MRD OUTPUT FILE PREPARATION AMPLIFICATION & MUITIPI FXING V1 2 SOFTWARE WITH VARIANT ON THE (ITD) & AMPLICONS ILLUMINA* CALL PURIFICATION MISEQ1

Figure 1: Workflow Summary

3.1. PCR Amplification of the Internal Tandem Duplication (ITD) Mutations of FLT3

FLT3 ITD or length mutations are caused by duplication and insertion of a portion of the *FLT3* gene that includes the region in and around the juxtamembrane (JM) region of the *FLT3* gene. These mutations vary in both the location and the length of the inserted duplicated DNA sequence. ITD mutations result in constitutive autophosphorylation and activation of *FLT3*.³

Primers are designed in and around the juxtamembrane region of the FLT3 gene.

3.2. Next-Generation Sequencing (NGS)

NGS technologies used in this assay rely on the amplification of genetic sequences using forward and reverse primers that include adapter and index tags. Amplicons generated with the *FLT3* ITD MRD Assay are quantified, pooled, and loaded onto a flow cell for sequencing with an Illumina® MiSeq™ sequencing platform. Specifically, the amplified products in the library are hybridized to oligonucleotides on a flow cell and are amplified to form local clonal colonies (bridge amplification). Four types of reversible terminator bases (RT-bases) are added and the sequencing strand of DNA is extended one nucleotide at a time. To record the incorporation of nucleotides, a CCD camera takes an image of the flourescence as each RT-base is added, and then cleaved to allow incorporation of the next base.

3.3. Sample Multiplexing

This product was designed to allow for two different levels of multiplexing in order to reduce costs and time for laboratories. The first level of multiplexing originates from the multiple indices that are provided with the assays. Each of the 24 dual-indexed master mixes acts as a unique barcode that allows amplicons from individual samples to be pooled together after PCR amplification to generate the sequencing library; the resulting sequences are sorted by the bioinformatic software, which identifies those that originated from an individual sample.

The second level of multiplexing originates from the ability to pool amplicons generated from multiple targets together, such as amplicon generated using Invivoscribe's *NPM1* MRD Assay (REF 14160019), to generate the library to be sequenced on a single flow cell (it is extremely important that each sample in the pool must have a unique index). When multiplexing amplicons of different gene targets it is important to use the appropriate sequencing chemistry. The number of sequencing cycles must be sufficient to sequence the largest amplicon in the multiplex. For example, when multiplexing a combination of *NPM1* MRD Assay and *FLT3* ITD MRD Assay amplicons together, use the MiSeq™ Reagent Kit v3 (600 cycle).

The number of samples that can be multiplexed onto a single flow cell is also dependent on the flow cell that is utilized. Illumina's standard flow cells (MiSeq™ Reagent Kit v3) can generate 22-25 million reads. To determine the number of reads per sample, divide the total number of reads for the flow cell by the number of samples that will be multiplexed.

3.4. Measurable Residual Disease Evaluation

The FLT3 ITD MRD is an NGS-based deep sequencing assay that can reliably identify DNA sequences specific to previously identified mutations at an allelic sensitivity level of 5×10^{-5} .

4. Reagents and Materials

NOTE: The *FLT3* ITD MRD Assay Kit, Table 1, is usable until the labeled kit expiration date when stored as described:

Reagents provided within this kit may incur up to 5 freeze-thaw cycles without measurable loss of performance. To reduce the number of freeze-thaws, aliquot the Positive and Negative Controls appropriately.

Table 1: Reagent List of the FLT3 ITD MRD Assay Kit (REF 14120019)

| Catalog Number | Reagent Name | Storage Temperature | Unit Quantity | Vials/ Kit |
|---------------------|----------------------|---------------------|---------------|------------|
| REF 24120249 | FLT3 ITD IA13 | | 75 μL/Vial | 1 |
| REF 24120259 | FLT3 ITD IA14 | | 75 μL/Vial | 1 |
| REF 24120269 | FLT3 ITD IA15 | | 75 μL/Vial | 1 |
| REF 24120279 | FLT3 ITD IA16 | | 75 μL/Vial | 1 |
| REF 24120289 | FLT3 ITD IA17 | | 75 μL/Vial | 1 |
| REF 24120299 | FLT3 ITD IA18 | | 75 μL/Vial | 1 |
| REF 24120309 | FLT3 ITD IA19 | | 75 μL/Vial | 1 |
| REF 24120319 | FLT3 ITD IA20 | | 75 μL/Vial | 1 |
| REF 24120329 | FLT3 ITD IA21 | | 75 μL/Vial | 1 |
| REF 24120339 | FLT3 ITD IA22 | | 75 μL/Vial | 1 |
| REF 24120349 | FLT3 ITD IA23 | | 75 μL/Vial | 1 |
| REF 24120359 | FLT3 ITD IA24 | ∫ | 75 μL/Vial | 1 |
| REF 24120369 | FLT3 ITD IB01 | | 75 μL/Vial | 1 |
| REF 24120379 | FLT3 ITD IB02 | -30°C → | 75 μL/Vial | 1 |
| REF 24120389 | FLT3 ITD IB03 | | 75 μL/Vial | 1 |
| REF 24120399 | FLT3 ITD IB04 | | 75 μL/Vial | 1 |
| REF 24120409 | FLT3 ITD IB05 | | 75 μL/Vial | 1 |
| REF 24120419 | FLT3 ITD IB06 | | 75 μL/Vial | 1 |
| REF 24120429 | FLT3 ITD IB07 | | 75 μL/Vial | 1 |
| REF 24120439 | FLT3 ITD IB08 | | 75 μL/Vial | 1 |
| REF 24120449 | FLT3 ITD IB09 | | 75 μL/Vial | 1 |
| REF 24120459 | FLT3 ITD IB10 | | 75 μL/Vial | 1 |
| REF 24120469 | FLT3 ITD IB11 | | 75 μL/Vial | 1 |
| REF 24120479 | FLT3 ITD IB12 | | 75 μL/Vial | 1 |
| REF 40883810 | FLT3 ITD MRD POS (+) | | 500 μL/Vial | 2 |
| REF 40920068 | FLT3 ITD MRD NEG (-) | | 500 μL/Vial | 2 |

Table 2: Additional Reagents, Materials, and Equipment Required (Not Provided)

| Reagent/Material | Suggested Reagents / Materials and Suppliers | Catalog No. | Notes |
|----------------------------------|------------------------------------------------------------|-------------|----------------------------|
| Taq Polymerase | Invivoscribe • FalconTaq™ DNA Polymerase | 60970130 | N/A |
| Molecular Biology Grade Water | N/A | N/A | Sterile, RNase/DNase -free |
| PCR Purification | Beckman Coulter • AMPure® XP Reagent for PCR Purification | N/A | N/A |
| Amplicon Quantification | N/A | N/A | N/A |
| Lab-grade Ethanol (200 Proof) | N/A | N/A | N/A |

Table 2: Additional Reagents, Materials, and Equipment Required (Not Provided)

| Reagent/Material | Suggested Reagents / Materials and Suppliers | Catalog No. | Notes |
|-----------------------------|----------------------------------------------------------------------------------------------------------------|-------------|----------------------------------------------------------------------------------|
| 1N NaOH N/A | | N/A | N/A |
| PhiX | Illumina® • PhiX Sequencing Control v3 | FC-110-3001 | N/A |
| Calibrated Pipettes | N/A | N/A | Must be able to accurately measure volumes between 0.5 μ L and 1000 μ L. |
| Filter Barrier Pipette Tips | N/A | N/A | Sterile, RNase/DNase/Pyrogen-free |
| PCR Plates or Tubes | N/A | N/A | Sterile, Skirted Plates |
| PCR Strip Caps or Foil Seal | N/A | N/A | N/A |
| DNA Low-Binding Tubes | N/A | N/A | DNA, DNase, RNase and PCR inhibitor free |
| Vortex Mixer | N/A | N/A | N/A |
| Microcentrifuge | N/A | N/A | N/A |
| Thermal Cycler | Thermo Fisher Scientific • Veriti™ 96-Well Fast Thermal Cycler | N/A | N/A |
| Magnetic Plate | Life Technologies ■ Magnetic Stand 96 | N/A | N/A |
| MiSeq™ | Illumina® • MiSeq™ Reagent Kit v3 (600-cycle)/ • MiSeq™ Reagent Kit v3 Box 1/ • MiSeq™ Reagent Kit v3 Box 2 | MS-102-3003 | Standard flow cell |
| MiSeq [™] Software | MiSeq™ Control Software (MCS) | N/A | v2.6.2.1 or later |

5. Software

5.1. FLT3 ITD MRD v1.2 Software

The *FLT3* ITD MRD v1.2 Software (**REF** 14120029) will be provided via a universal serial bus (USB) where the customer will download and install the software in their local hardware infrastructure. The customer is provided the download and end-user instructions for installation and execution of software.

5.1.1. Required Equipment

- 5.1.1.1. A Linux server with the following minimum requirements:
 - Processor: Minimum of 16 cores required
 - Hard Drive: At least 50 GB of free disk space
 - RAM: 64 GB required; 128 GB or more recommended
 - Operating System: Ubuntu release 18.04+
 - All FLT3 ITD MRD v1.2 Software dependencies must be installed:
 - Docker installed on host machine

6. Warnings and Precautions

Please read the Instructions for Use carefully prior to starting the assay procedure and follow each step closely.

- RUO This product is for Research Use Only.
- The assay is only validated for use on the MiSeq[™] with MCS. The assay must be used as a system. Do not substitute other manufacturers' reagents.
- Dilution, reducing amplification reaction volumes, or other deviation in this protocol may affect the performance of this test and/or nullify any limited sublicense that comes with the purchase of this testing kit.
- Do not mix or combine reagents from kits with different lot numbers.
- Materials are stable until the labeled expiration date when stored and handled as directed. Do not use kits beyond their expiration date.
- Dispose of unused reagents and waste in accordance with country, federal, state and local regulations.
- Track the number of freeze thaw cycles.
- Perform all laboratory procedures with standard personal protective equipment (gloves, laboratory coats, and protective eye wear). Follow good laboratory practices and universal precautions when working with specimens. Do not pipette by mouth. Do not eat, drink, or smoke in laboratory work areas. Wash hands thoroughly after handling specimens and assay reagents. Handle specimens in approved biological safety containment facilities and open only in certified biological safety cabinets.
- Due to the analytical sensitivity of this test, use extreme care to avoid the contamination of reagents or amplification mixtures with samples, controls, or amplified materials. Use fresh, aerosol-resistant pipette tips between samples and between dispensing reagents. Closely monitor all reagents for signs of contamination (e.g., negative controls giving positive signals). Discard reagents suspected of contamination.
- To minimize contamination, wear clean gloves when handling samples and reagents and routinely clean work areas and pipettes prior to performing PCR.
- Autoclaving does not eliminate DNA contamination. Follow uni-directional work flow in the PCR laboratory between separate work areas; begin with specimen preparation, then to amplification, and finally to sequencing. Do not bring amplified DNA into the areas designated for specimen preparation.
- Dedicate all pipettes, pipette tips, and any equipment used in a particular area to that area of the laboratory.
- Use sterile, disposable plastic ware whenever possible to avoid RNase, DNase, or cross-contamination.
- All instruments and equipment must be maintained and calibrated per the manufacturers' recommendations.

6.1. Cyber Security Precautions

- Computers and networks are susceptible to security risk if not secured and actively updated. Proper computer
 and network security help ensure data is not compromised, lost, or damaged due to preventable cyber risks. Equip
 all computers with up to date and active antivirus software.
- Filter and secure network traffic with a firewall.
- Keep data on local computers to reduce cyber security risks that may be present in transferring sensitive data over a network.
- Install software only for the local user to prevent unauthorized use of the software.
- Ensure Windows and Adobe Acrobat Reader are always updated to the latest available security patches.
- Ensure the default PDF reader in Windows is set to Adobe Acrobat Reader. Opening sample and run reports in an internet browser may lead to cybersecurity risks of subject data.

7. Specimen Collection and Preparation

7.1. Precautions

 Biological specimens from humans may contain potentially infectious materials. Handle all specimens according to your institute's Bloodborne Pathogen program and/or Biosafety Level 2.

7.2. PCR-Interfering Substances

- Divalent cation chelators
- Low retention pipette tips
- EDTA (not significant at low concentrations)

7.3. Specimen Requirements and Stability

- This assay tests genomic DNA (DNA) from the following sources:
 - Peripheral blood or bone marrow in EDTA or sodium heparin
- Specimens can be stored at 4°C or at Room Temperature for up to 7 days after the collection date prior to DNA isolation.

7.4. Sample Preparation

- 7.4.1. Extract DNA from peripheral blood or bone marrow within 7 days of collection.
- 7.4.2. Quantify DNA samples using a fluorescence-based method specific to double-stranded DNA (dsDNA).

NOTE: Use a quantification method appropriate for dsDNA. If DNA concentration is <20 ng/ μ L, refer to troubleshooting guidelines in section 10. Troubleshooting.

7.4.3. Dilute an aliquot of the stock DNA to 20 $ng/\mu L$ in molecular grade water using low-binding surface tubes using Equation 1.

Equation 1: Equation to Normalize DNA
$$V_i = rac{\left(V_f \, x \, 20 \, rac{ng}{\mu L}
ight)}{C_i}$$

- **C**_i = DNA concentration from microvolume fluorometer reading
- V_i = volume of undiluted DNA to dilute
- V_f = final volume of diluted DNA
- $V_f V_i$ = amount of molecular grade water to add to V_i

8. Assay Procedure

8.1. PCR Setup

Perform all thermal cycler steps including installation, operation, calibration, cleaning and maintenance procedures according to the manufacturer's instructions unless stated otherwise below.

NOTE: Minimize the amount of time Taq DNA Polymerase is out of -30°C to -15°C storage.

- A Positive Control, Negative Control, and No Template Control must be included in each run. Up to 21 samples (in singlicate) may be included in one run.
- The No Template Control must be run with each PCR step using molecular grade water to verify absence of amplicon contamination.
 - 8.1.1. Remove Control tubes (*FLT3* ITD MRD POS (+)and *FLT3* ITD MRD NEG (-)) from appropriate storage and thaw at room temperature.
 - 8.1.2. Pipette 35 μ L of normalized sample DNA and Controls at 20 ng/ μ L to a 96-well PCR plate in a staggered layout, as shown in Figure 2. For No Template Control, pipette 35 μ L of molecular biology grade water.

| | | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|----------------------|---|----------------------|---------------|---|---|---|---|---|---|---|----|----|----|
| Key | Α | <i>FLT3</i> ITD IA13 | | | | | | | | | | | |
| Sample | В | | FLT3 ITD IA17 | | | | | | | | | | |
| FLT3 ITD MRD POS (+) | | FLT3 ITD IA14 | | | | | | | | | | | |
| FLT3 ITD MRD NEG (-) | D | | | | | | | | | | | | |
| NTC | E | FLT3 ITD IA15 | | | | | | | | | | | |
| | F | | | | | | | | | | | | |
| | G | <i>FLT3</i> ITD IA16 | | | | | | | | | | | |
| | н | | | | | | | | | | | | |

Figure 2: Example of Staggered Samples

- 8.1.3. Allow the ITD Master Mixes to thaw; then gently vortex to mix and centrifuge.
- 8.1.4. Perform the remaining PCR plate setup on a cold block.
- 8.1.5. Cover all of the wells except the working column, i.e. column 1, with a foil seal.
 - To prevent accidental cross contamination of the indexes, the remaining PCR reagents must be added one column at a time, one reagent at a time. All reagents will be added per column, and the column sealed before moving onto the next.
- 8.1.6. Opening only one Master Mix at a time, for each exposed PCR reaction, pipette 15 μ L of the respective PCR Master Mix.
 - This will yield a total volume of 50 μL per well. Use a Master Mix with a unique sequencing index for each DNA sample or Control in the run.

NOTE: It is very important to only have one Master Mix tube open at a time to prevent cross contamination.

- 8.1.7. Pipette 1 µL of Taq DNA Polymerase to each exposed well.
- 8.1.8. Cap the working column with 8-well strip caps, ensuring a tight seal over each well.
- 8.1.9. Repeat starting from section 8.1.5 for the remaining plate columns.
- 8.1.10. Gently vortex the capped PCR plate and quickly centrifuge to gather all droplets to the bottom of the wells.
 - When vortexing try to keep the reaction in the bottom of the tube and off the strip caps.
- 8.1.11. Place the PCR plate in a pre-heated thermal cycler and close the lid.
- 8.1.12. Amplify the PCR plate according to the PCR program in Table 3.
 - If not immediately continuing onto the next steps, the PCR product plate may be stored at 4°C for up to 72 hours.

Table 3: PCR Amplification Thermal Cycler Program

| Step | Temperature | Time | Cycles | | |
|-------------------------|-------------|------------|--------|--|--|
| 1 | 95°C | 7 minutes | 1 | | |
| 2 | 95°C | 45 seconds | | | |
| 3 | 60°C | 45 seconds | 25x | | |
| 4 | 72°C | 90 seconds | | | |
| 5 | 72°C | 10 minutes | 1 | | |
| 6 | 4°C | ∞ | Hold | | |
| Ramp rate: 2.9°C/second | | | | | |

NOTE: Set the heated lid to 105°C and the reaction volume to 50 µL

8.2. First PCR Product Purification (0.8x Cleanup)

NOTE: AMPure® XP Reagent for PCR purification is viscous and needs to be pipetted slowly. Ensure beads are homogenized prior to pipetting.

- 8.2.1. Remove AMPure® XP Reagent from storage and allow to equilibrate to room temperature for at least 30 minutes before use.
- 8.2.2. Transfer the appropriate volume of AMPure® XP Reagent needed to a new reservoir to minimize the risk of contamination by pipette tips.
 - 8.2.2.1. The required volume of AMPure[®] XP Reagent = $(n + 2) \times 40 \mu L$ (n is number of samples/controls to be purified).
- 8.2.3. Prepare 80% ethanol using 100% ethanol and molecular biology grade water (0.5 mL for each sample/control to be purified).
- 8.2.4. Remove the PCR product plate from the thermal cycler or 4°C storage and centrifuge to ensure all liquid is at the bottom of the wells.
- 8.2.5. Pipette 40 μL of the aliquoted, room temperature AMPure® XP Reagent to each PCR product to be purified. Mix by pipetting up and down 10 times. The color of the mixture should appear homogenous after mixing.
- 8.2.6. Incubate 5 minutes at room temperature.
- 8.2.7. Place the mixed samples/controls on a 96-well plate magnet and allow the magnetic particles to separate for 2 minutes.
 - Keep the plate on the magnetic stand at all times during this procedure, until step 8.2.15.
- 8.2.8. Aspirate 85 μ L of the clear supernatant and discard.
 - Avoid removing any magnetic particles.
- 8.2.9. Keeping the plate on the magnetic stand, add 200 μL of 80% ethanol to each sample/control well without mixing.
- 8.2.10. Incubate at room temperature on the magnetic stand for 30 seconds.
- 8.2.11. Aspirate 200 μ L of the ethanol and discard.
- 8.2.12. Repeat steps 8.2.9 through 8.2.11, for a total of 2 washes.
- 8.2.13. Using a fine-tipped pipette, aspirate and discard any excess ethanol.
- 8.2.14. With the plate still on the magnet stand, allow the magnetic particles to air-dry for 5 minutes.
- 8.2.15. Remove the plate from the magnet stand and add 50 µL of 10 mM Tris-HCl, pH 8.0 buffer.
 - Mix by pipetting until homogeneous.
 - Ensure all magnetic particles are in solution.
- 8.2.16. Incubate at room temperature for 2 minutes.
- 8.2.17. Place the plate on the magnet stand for 2 minutes or until the supernatant has cleared.
- 8.2.18. Transfer 47 μL of eluate to a fresh plate. If continuing onto the next section, mix by pipetting 3 times.
- 8.2.19. Centrifuge to ensure all of the solution is at the bottom of the wells.

• If not immediately continuing to the next step, the purified amplicons can be stored at 4°C for up to 72 hours or at -20°C for up to 2 weeks.

8.3. Second PCR Product Purification (0.7x Cleanup)

- 8.3.1. If not already at room temperature, remove AMPure® XP Reagent from storage and allow to equilibrate to room temperature for at least 30 minutes before use.
- 8.3.2. Transfer the appropriate volume of AMPure® XP Reagent needed to a new reservoir to minimize the risk of contamination by pipette tips.
 - 8.3.2.1. The required volume of AMPure® XP Reagent = $(n + 2) \times 32.9 \mu L$ (n is number of samples/controls to be purified).
- 8.3.3. Prepare 80% ethanol using 100% ethanol and molecular biology grade water (0.5 mL for each sample/control to be purified).
- 8.3.4. If purified amplicon was stored at -20°C, remove from storage and thaw. Centrifuge the PCR product plate.
- 8.3.5. Pipette 32.9 μL of the aliquoted, room temperature AMPure® XP Reagent to each PCR product to be purified. Mix by pipetting up and down 10 times. The color of the mixture should appear homogenous after mixing.
- 8.3.6. Incubate 5 minutes at room temperature.
- 8.3.7. Place the mixed samples/controls on a 96-well plate magnet and allow the magnetic particles to separate for 2 minutes.
 - Keep the plate on the magnetic stand at all times during this procedure, until step 8.3.15.
- 8.3.8. Aspirate 75 μ L of the clear supernatant and discard.
 - Avoid removing any magnetic particles.
- 8.3.9. Keeping the plate on the magnetic stand, add 200 μL of 80% ethanol to each sample/control well without mixing.
- 8.3.10. Incubate at room temperature on the magnetic stand for 30 seconds.
- 8.3.11. Aspirate 200 µL of the ethanol and discard.
- 8.3.12. Repeat steps 8.3.9 through 8.3.11, for a total of 2 washes.
- 8.3.13. Using a fine-tipped pipette, aspirate and discard any excess ethanol.
- 8.3.14. With the plate still on the magnet stand, allow the magnetic particles to air-dry for 5 minutes.
- 8.3.15. Remove the plate from the magnet stand and add 50 µL of 10 mM Tris-HCl, pH 8.0 buffer.
 - Mix by pipetting until homogeneous.
 - Ensure all magnetic particles are in solution.
- 8.3.16. Incubate at room temperature for 2 minutes.
- 8.3.17. Place the plate on the magnet stand for 2 minutes or until the supernatant has cleared.
- 8.3.18. Transfer 47 μ L of eluate to a fresh plate. Mix by pipetting 3 times.
- 8.3.19. Centrifuge to ensure all of the solution is at the bottom of the wells.
 - If not immediately continuing to the next step, the purified amplicons can be stored at 4°C for up to 72 hours or at -20°C for up to 2 weeks.

8.4. Quantification and Validity of Purified PCR Products

Use an appropriate method that assesses both concentration and sizing of purified dsDNA PCR products between 200 bp - 1000 bp. Perform all quantification steps including installation, operation, calibration, cleaning and maintenance procedures according to the manufacturer's instructions.

8.4.1. Verify control and sample validity per Table 4.

Table 4: Control and Sample Quantification Validity

| Validity Requirement | Acceptance Criteria |
|----------------------------------------------------|---------------------|
| FLT3 ITD MRD POS (+), FLT3 ITD MRD NEG (-), Sample | ≥1.00 nM |
| No Template Control | ≤1.00 nM |

- 8.4.2. If a control or sample molarity (nM) value does not meet the acceptance criteria, refer to section 10. Troubleshooting.
- 8.4.3. If not immediately continuing to the next step, the amplicons can be stored at 4°C for up to 2 weeks.

8.5. Pooling and Quantification of the Library

Accurate quantification and dilution of the pool loaded onto the MiSeq[™] flow cell is critical for generating optimal cluster density on the flow cell and obtaining high-quality data in a sequencing run.

- 8.5.1. Determine the desired library pool's target molarity (nM) value between 2-10 nM.
- 8.5.2. Normalize the Positive Control, Negative Control, and sample to the target molarity (nM) value using 10 mM Tris-HCl, pH 8.0.
- 8.5.3. Combine equal volumes of the normalized controls and sample in a volume appropriate tube. Include an equal volume of No Template Control.
- 8.5.4. Gently vortex the tube to thoroughly mix and quickly centrifuge.
- 8.5.5. Quantify the library pool using a method that is appropriate for both concentration and sizing of purified dsDNA PCR products between 200 bp 1000 bp.
- 8.5.6. If not immediately continuing to the next step, the library pool can be stored at -20°C. Avoid unnecessary freezethaw cycles.

8.6. Preparation of the Library Pool for the MiSeq™ Sequencing Run

The concentration of the final library pool to be loaded onto the MiSeq™ Reagent v3 Cartridge is 14 pM.

- 8.6.1. If stored at -20°C, thaw library pool. Gently vortex the tube to thoroughly mix and quickly centrifuge.
- 8.6.2. Create a 20 µL denatured library pool at a target concentration of 1.75 nM by first calculating the volume of final library, X, using information in Table 5 and Equation 2. Use the final library molarity (nM) value determined in section 8.5.5 as the denominator in Equation 2.

Table 5: Library Denaturation

| Reagent | Volume (μL) |
|-------------------------------|-----------------------|
| Final Library | X (use Equation 2) |
| Diluent (10mM Tris-HCl, pH 8) | Y (use Equation 3) |
| 10 nM PhiX (12.5%) | 0.5 |
| 1N NaOH | 2 |
| Total Reaction Volume | 20 |

$$X = \frac{\left(Target \ nM \ x \ Total \ Reaction \ Volume \ (\mu L)\right)}{Final \ Library \ nM}$$

$$X = \frac{1.75 \ nM \ x \ 20 \ \mu L}{Final \ Library \ nM}$$

8.6.3. Second, calculate the volume of diluent, Y, to create a 20 μL denatured library pool at 1.75 nM using information in Table 5 and Equation 3.

Equation 3: Equation to Calculate Diluent

$$Y = Total \text{ Reaction } Volume \ (\mu L) - Volume \ of \ PhiX \ (\mu L) - Volume \ of \ NaOH \ (\mu L) - Volume \ of \ Final \ Library \ (\mu L)$$

$$Y = 20 \ \mu L - 0.5 \ \mu L - 2.0 \ \mu L - X \ \mu L$$

- 8.6.4. Add the calculated volume of diluent (Y) into a 1.5 mL low binding tube.
- 8.6.5. Add the calculated volume of library (*X*) into the tube.
- 8.6.6. Add 0.5 μ L 10 nM PhiX into the tube.
- 8.6.7. Add 2 μL of 1N NaOH into the tube. Mix well by flicking the tube 5 times. Briefly centrifuge the tube.
- 8.6.8. Incubate the tube at room temperature for 5 minutes to denature the double-stranded DNA into single-stranded DNA.
- 8.6.9. Immediately upon the completion of the 5-minute incubation, add 980 μ L of pre-chilled HT1 buffer to the reaction.
- 8.6.10. Vortex the tube to thoroughly mix and quickly centrifuge. The denatured Library Pool reaction is now at 40 pM (Library Pool pM + PhiX pM).
- 8.6.11. Combine 350 μ L of the 40 pM denatured Library Pool and 650 μ L of pre-chilled HT1 to create a Final Library Pool at 14 pM. Vortex the tube to thoroughly mix and quickly centrifuge.

8.7. Preparation of the MiSeq™ Reagent Cartridge

Prepare reagent cartridge according to the following sections of the MiSeq System Guide (EEE 15027617).

- Thaw Reagent Cartridge
- Inspect the Reagent Cartridge

8.8. MiSeg™ Sample Sheet Creation

The MiSeq[™] sample sheet will be provided via USB (REF 14120029).

NOTE: Do not make any modifications to the sample sheet other than those instructed below. Doing so may inhibit analysis.

- 8.8.1. Using the provided sample sheet, ensure the following parameters are correct:
 - Workflow: GenerateFASTQApplication: FASTQ Only
 - Assay: NexteraChemistry: AmpliconReads 1 and 2: 301
- 8.8.2. Do not edit or delete lines 22 and 23 on the provided sample sheet.
- 8.8.3. Use the provided sample sheet and follow the sample character and format rules to input Sample_ID, Sample_Name, I7_Index_ID, index, I5_index_ID, index2, and a Description beginning on line 24:
 - Sample ID and Sample Name column requirements:
 - Underscores and blank spaces are not allowed; use a dash "-" if needed. Use only these characters: A/a through Z/z, 0 through 9, -
 - Use the Master Mix ID or sample index, e.g. IA13
 - Sample_ID and Sample_Name must be identical
 - o Example: FLT3-ITD-MRD-IA13-Pos
 - *I7_Index_ID* and *I5_Index_ID* column requirements:
 - Correlate I7_Index_ID and I5_index_ID to the Master Mix ID, e.g., if Master Mix ID is IA13, input N701 and N502, per Table 6
 - o Index ID pairs must be unique and not shared across multiple samples on the same sequencing run
 - index and index2 column requirements:
 - o Enter a unique index sequence
 - Index sequence pairs must be unique and not shared across multiple samples on the same sequencing run
 - Description column requirements:
 - Must contain DNA amount in ng, e.g. 700ngInput. It is necessary to include "ng" with the numerical value and exclude space between "ng" and "Input."

Table 6: Indexes

| Master Mix ID | I7_Index_ID | I7_Index Sequence | I5_index_ID | I5_index Sequence |
|---------------|-------------|-------------------|-------------|-------------------|
| IA13 | N701 | TAAGGCGA | N502 | CTCTCTAT |
| IA14 | N702 | CGTACTAG | N504 | AGAGTAGA |
| IA15 | N703 | AGGCAGAA | N506 | ACTGCATA |
| IA16 | N704 | TCCTGAGC | N508 | CTAAGCCT |
| IA17 | N705 | GGACTCCT | N501 | TAGATCGC |
| IA18 | N706 | TAGGCATG | N503 | TATCCTCT |
| IA19 | N707 | CTCTCTAC | N505 | GTAAGGAG |
| IA20 | N708 | CAGAGAGG | N507 | AAGGAGTA |
| IA21 | N709 | GCTACGCT | N502 | CTCTCTAT |
| IA22 | N710 | CGAGGCTG | N504 | AGAGTAGA |
| IA23 | N711 | AAGAGGCA | N506 | ACTGCATA |
| IA24 | N712 | GTAGAGGA | N508 | CTAAGCCT |
| IB01 | N702 | CGTACTAG | N501 | TAGATCGC |
| IB02 | N703 | AGGCAGAA | N503 | TATCCTCT |
| IB03 | N704 | TCCTGAGC | N505 | GTAAGGAG |
| IB04 | N705 | GGACTCCT | N507 | AAGGAGTA |

Table 6: Indexes

| Master Mix ID | I7_Index_ID | I7_Index Sequence | I5_index_ID | I5_index Sequence |
|---------------|-------------|-------------------|-------------|-------------------|
| IB05 | N706 | TAGGCATG | N502 | CTCTCTAT |
| IB06 | N707 | CTCTCTAC | N504 | AGAGTAGA |
| IB07 | N708 | CAGAGAGG | N506 | ACTGCATA |
| IB08 | N709 | GCTACGCT | N508 | CTAAGCCT |
| IB09 | N710 | CGAGGCTG | N501 | TAGATCGC |
| IB10 | N711 | AAGAGGCA | N503 | TATCCTCT |
| IB11 | N712 | GTAGAGGA | N505 | GTAAGGAG |
| IB12 | N701 | TAAGGCGA | N507 | AAGGAGTA |

8.8.4. Save the CSV file, explicitly named "SampleSheet.csv," to a folder accessible by the MiSeq™ instrument.

8.9. MiSeq™ Reagent Cartridge Loading

Load 600 μ L of the 14 pM Final Library Pool onto a MiSeq[™] Reagent v3 Cartridge according to the following sections of the *MiSeq System Guide* (EEF 15027617 v05).

Load Sample Libraries

8.10. MiSeq™ Run Start

- 8.10.1. Start the MiSeq[™] run according to the *MiSeq System Guide* (REF 15027617).
- 8.10.2. When selecting the run, review run parameters and verify the cycling conditions.

NOTE: To reduce the occurrence of run-to-run cross-contamination, it is recommended to alternate sequencing runs with IA or IB master mixes, or alternatively, run a non-FLT3 ITD MRD Assay run between FLT3 ITD MRD Assay sequencing runs.

8.11. Acceptance Criteria

8.11.1. Verify sequencing run validity in Table 7.

Table 7: Run Validity

| Validity Requirement | Acceptance Criteria |
|------------------------------------|---------------------------------------------------------------------------------|
| MiSeq [™] Cluster Density | ≥500 K/mm² |
| Total MiSeq™ Run Q30 Score | ≥60% |
| MiSeq™ Reads Pass Filter | ≥10 Million |
| FLT3 ITD MRD POS (+) | Determined to be "Detected" with a 30 bp ITD and a VRF of ≥1.2x10 ⁻⁷ |
| FLT3 ITD MRD NEG (-) | "Not Detected" for an ITD |
| No Template Control | ≤1,000 reads and is determined to be "Undetermined" |

8.11.2. If the sequencing run does not meet all acceptance criteria, see section 10. Troubleshooting.

9. Data Analysis

9.1. FLT3 ITD MRD v1.2 Software

NOTE: If using Windows 10, the formatting of the MiSeq™ samplesheet.csv output may be impacted by the MiSeq Control Software. When executing software analysis, ensure proper formatting of the SampleSheet.csv by using the sample sheet created in section 8.8.

- 9.1.1. Use the FLT3 ITD MRD v1.2 Software (EEF 14120029) for data analysis.
- 9.1.2. Refer to the following README files included in the software package to understand software execution via command-line or via the "DREW" rest-API interface:
 - 9.1.2.1. DREW README.md
 - 9.1.2.2. Distributable_README.txt

9.2. Sample Interpretation

- 9.2.1. A minimum of 3 variant supporting reads is necessary for a positive call.
- 9.2.2. Minimum reads required for 95% confidence in negative call at 5x10⁻⁵, given 700 ng DNA input, is 157,600.
 - 9.2.2.1. A negative sample with fewer than 157,600 reads is reported as Undetermined.

9.3. ITD and Insertion Variant Detection

- 9.3.1. The observed variant read frequency (VRF) is provided in the "ITD_VRF" column.
- 9.3.2. The number of variant-supporting reads is provided in the "ITD READS" column.
- 9.3.3. The software can detect ITDs from 9 base pairs up to a length of 252 base pairs and insertions from 3 base pairs to less than the maximum ITD size, depending on the location of the duplication. If the location of the duplication is close to the end of exon 14, the detection length is limited to 180 base pairs.
- 9.3.4. In the TSV output file, an asterisk (*) character next to ITD_Size indicates the variant is identified as a simple insertion and not a complex tandem duplication insertion variant.

9.4. Sequence Annotations

- 9.4.1. Indel sequence information can be found following this file path: <ANALYSIS_DIR>/<subdir>/raw_results/indels/<sampleName>.insertion.out.
- 9.4.2. Match the insertion from the final output TSV file according to the insertion size.
- 9.4.3. Chromosomal location is not provided.

10. Troubleshooting

Table 8: Troubleshooting Guide

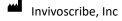
| General Failure Category | Description of Failure | Action |
|-----------------------------------|-------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Sample and Reagent Preparation | Sample DNA quantity is < 20 ng/μL by a fluorescence- based method | Reprocess the primary specimen (peripheral blood) one time (if within the validated stability claim. If outside the stability claim, request an additional specimen collection. |
| | Constitution of NTC in 1 00 and | 1. If the NTC is between 1.00 nM – 2.00 nM, manually check to confirm no peak is present between 200 – 1000 bp range to pass. |
| | Concentration of NTC is >1.00 nM | 2. If NTC >2.00nM, sample contamination is likely; therefore, identify if there is enough starting material to re-amplify <u>all</u> samples. If so, repeat protocol from the beginning. |
| Quantification of Sample | If undiluted amplicon is <1.00 nM | Re-quantify with a new dilution. If a greater value is obtained, continue with corrected value. If same value is obtained, re-purify and repeat the protocol from the beginning for that sample. After re-amplification, it may be pooled with the amplicons from the first run. If there is not enough sample to repeat amplification, continue through the end of the protocol pooling the max volume (10-40 μL) of sample. |
| MiSeq™ Run | MiSeq™ Run Q30 is <60% | 1. Re-quantify the pooled library and re-sequence. |
| MiSeq™ Run | MiSeq™ Reads Pass Filter is <10 Million | 1. Re-quantify the pooled library and re-sequence. |

11. References

- 1. Daver, N. et al., (2019). <u>Targeting FLT3 mutations in AML: review of current knowledge and evidence</u>. *Leukemia*. 33, 299–312.
- 2. Krönke, J. et al., (2013). Clonal evolution in relapsed NPM1-mutated acute myeloid leukemia. Blood 122(1), 100-8.
- 3. Murphy, KM. et al., (2003). <u>Detection of FLT3 Internal Tandem Duplication and D835 Mutations by a Multiplex Polymerase Change Reaction and Capillary Electrophoresis Assay</u>. *Journal of Molecular Diagnostics* 5, 96-102.
- 4. Levis, MJ. et al., (2018). A next-generation sequencing—based assay for minimal residual disease assessment in AML patients with FLT3-ITD mutations. Blood Adv. 2(8), 825-831.
- MiSeq System Guide (REF 15027617).
- http://www.illumina.com

12. Technical and Customer Service

Contact Information



10222 Barnes Canyon Road | Building 1 | San Diego | California 92121-2711 | USA

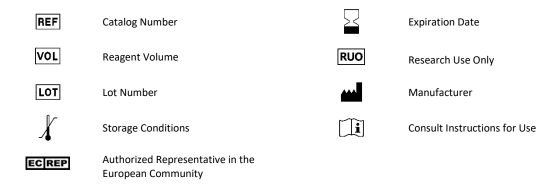
Phone: +1 858 224-6600 | Fax: +1 858 224-6601 | Business Hours: 7:00AM - 5:00 PM PST/PDT

Technical Service: support@invivoscribe.com | Customer Service: sales@invivoscribe.com | Website: www.invivoscribe.com | Customer Service: sales@invivoscribe.com | Website: www.invivoscribe.com | Webs

Technical and Customer Service Representatives are available Monday through Friday to answer phone, e-mail, or website inquiries.

13. Symbols

The following symbols are used in Invivoscribe NGS product labeling.



14. Legal Notice

Invivoscribe, Inc. (Invivoscribe®) is committed to providing the highest quality products. Invivoscribe® warrants that the products meet or exceed the performance standards described in the Instructions For Use, as to products with such an insert. If a product is covered by product specifications and does not perform as specified, our policy is to replace the product or credit the full purchase price. No other warranties of any kind, expressed or implied, are provided by Invivoscribe®. Invivoscribe® liability shall not exceed the purchase price of the product. Invivoscribe® shall have no liability for direct, indirect, consequential or incidental damages arising from the use, results of use, or inability to use its products; product efficacy under purchaser controlled conditions in purchaser's laboratory must be established and continually monitored through purchaser defined and controlled processes including but not limited to testing of positive, negative, and blank controls every time a sample is tested. Ordering, acceptance, and use of product constitutes purchaser acceptance of sole responsibility for assuring product efficacy and purchaser agreement to the limitation of liability set forth in this paragraph.

Many of these products require nucleic acid amplification methods such as Polymerase Chain Reaction (PCR). No license under these patents to use amplification processes or enzymes is conveyed expressly or by implication to the purchaser by the purchase of this product.

© 2023 Invivoscribe, Inc. All rights reserved. The trademarks mentioned herein are the property of Invivoscribe, Inc. and/or its affiliates, or (as to the trademarks of others used herein) their respective owners.

This product is for Research Use Only; not for use in diagnostic procedures.