Intended Use

Myriad MyChoice® CDx is a next generation sequencing-based *in vitro* diagnostic test that assesses the qualitative detection and classification of single nucleotide variants, insertions and deletions, and large rearrangement variants in protein coding regions and intron/exon boundaries of the *BRCA1* and *BRCA2* genes and the determination of Genomic Instability Score (GIS), which is an algorithmic measurement of Loss of Heterozygosity (LOH), Telomeric Allelic Imbalance (TAI), and Large-scale State Transitions (LST) using DNA isolated from formalin-fixed paraffin embedded (FFPE) tumor tissue specimens.

The results of the test are used as an aid in identifying ovarian cancer patients with positive homologous recombination deficiency (HRD) status, who are eligible, because of a positive test result for deleterious or suspected deleterious mutations in *BRCA1* or *BRCA2* genes, or may become eligible, because of a positive test result for deleterious or suspected deleterious mutations in *BRCA1* or *BRCA2* genes or a positive Genomic Instability Score, for treatment with the targeted therapy listed in Table 1 in accordance with the most recently approved therapeutic product labeling.

### Table 1: Companion diagnostic indications

<table>
<thead>
<tr>
<th>Tumor Type</th>
<th>Biomarker</th>
<th>Therapy</th>
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<tbody>
<tr>
<td>Ovarian Cancer</td>
<td>Myriad HRD, defined as:</td>
<td>Lynparza® (olaparib)*</td>
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<td></td>
<td>• deleterious or suspected deleterious mutations in <em>BRCA1</em> and <em>BRCA2</em> genes and/or</td>
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<td></td>
<td>• positive Genomic Instability Score</td>
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* Refer to the drug label for HRD definition for olaparib monotherapy or combination therapy.

Detection of deleterious or suspected deleterious *BRCA1* and *BRCA2* mutations and/or positive Genomic Instability Score in ovarian cancer patients is also associated with enhanced progression-free survival (PFS) from Zejula® (niraparib) maintenance therapy in accordance with the most recently approved therapeutic product labeling. This assay is for professional use only and is to be performed only at Myriad Genetic Laboratories, Inc., a single laboratory site located at 320 Wakara Way, Salt Lake City, UT 84108.
MyChoice® CDx Technical Information
www.myriad.com/technical-specifications

Contraindication

- There are no known contraindications.

Warnings and Precautions

- There are no known warnings or precautions.

Limitations

- For in vitro diagnostic use
- For professional use only
- For prescription use only
- This test identifies germline and somatic variants in the tumor but does not distinguish between the two.
- Reduced hybridization efficiency of DNA fragments spanning long insertions and deletions (indels), or rearrangements may result in under-representation of mutant DNA molecules in the final sequencing library. This will result in a reduction in the observed frequency of sequence reads spanning the mutation.
- Indels >25 bp in length can be detected by this assay. However, the ability to detect any particular indel may be impacted by the location and nature of the mutation, the local sequence context, the DNA quality, and the non-tumor DNA content in the sample provided.
- This test has been designed to detect genomic rearrangements including large rearrangements (LRs) involving the promoter and coding exons of BRCA1 and BRCA2, however, the detection of large rearrangement deletions and duplications is dependent on the quality of the submitted specimen.
- Whole gene duplications and deletions may not be detected by the MyChoice CDx assay.
- Other terminal duplications are reported as variants of uncertain significance.
- This analysis is believed to rule out the majority of abnormalities in the genes analyzed. There may be uncommon genetic abnormalities such as specific insertions, inversions, and certain regulatory mutations that will not be detected by the MyChoice CDx assay.
- Samples with DNA input of <30ng may exhibit reduced read coverage, potentially affecting the assay’s sensitivity in detecting variants with low allele frequencies (<10%). In addition, samples with a low DNA input (< 30ng) are more likely to result in a failure of GIS analysis or to detect large rearrangements.
- Alterations at allele frequencies below the established limit of detection may not be detected consistently.

Test Principle

The MyChoice CDx test determines a patient’s Myriad HRD Status by detecting single nucleotide variants (SNVs), variants in homopolymer stretches, indels, and LRs in the BRCA1 and BRCA2 genes and determining a GIS using DNA obtained from FFPE ovarian tumor tissue. A positive Myriad HRD Status result is due to either the presence of a pathogenic mutation in BRCA1 and/or BRCA2 (sequencing and/or LR) [positive Tumor Mutation BRCA1/BRCA2 Status (tBRCA1/2 Status)] and/or a positive Genomic Instability Score (i.e., GIS above a defined threshold [positive GIS Status]).
The assay employs a single DNA extraction method from FFPE specimens, 7.5-200 ng of which undergoes multiple steps including fragmentation, end repair and adenylation, adapter ligation, library construction/amplification, hybridization and capture, sequencing, and data analysis.

The MyChoice CDx test is composed of the following major processes:

- Tumor Sample Collection and Shipping
- Tumor genomic DNA Extraction
- DNA Processing using the following assays:
  - MyChoice CDx next generation sequencing is used to detect sequence variants and genomic rearrangements (i.e., large deletions and duplications) in BRCA1 or BRCA2, and genomic instability analysis.
- BRCA1 and BRCA2 Variant Classification
- Genomic Instability Score (GIS) Calculation
- Results Reporting

Summary and Explanation

The MyChoice CDx device is a companion diagnostic for Lynparza® (olaparib) a poly ADP-ribose polymerase (PARP) inhibitor. Patients whose tumors have a positive GIS Status and/or pathogenic variants in BRCA1 or BRCA2 show improved progression free survival when treated with Zejula® (niraparib) compared to placebo in the treatment setting.

Test Kit Contents

A sample collection kit provided by Myriad is used by the ordering laboratories/physicians. The collection kits contain the following components:

- Slide Container
- Tumor Block Container
- Collection Instructions
- Mailing Instructions
- Ice Pack

Instruments

The MyChoice CDx device is intended to be performed with the following instruments, as identified by specific serial numbers:

- Eppendorf MasterCycler PRO-S 96 well Thermocycler
- Illumina NovaSeq 6000 Next Generation Sequencer

Sample Collection and Test Ordering

To order MyChoice CDx testing, the Test Request Form (TRF) included in the test kit must be fully completed. Please refer to the MyChoice CDx Collection Instructions and Mailing Instructions for further details about collecting formalin-fixed paraffin-embedded (FFPE) tumor samples and mailing the samples to Myriad.
Test Results and Interpretation

Patients evaluated with the MyChoice CDx test that have clinically relevant tumor genomic instability and/or have been determined to carry a deleterious or suspected deleterious BRCA1 or BRCA2 mutation can be considered for treatment with Lynparza® (olaparib) under the supervision of a physician.

Upon completion of testing at Myriad, a test report will be sent to the designated physician. The results of each test component, along with the interpretation of any reportable BRCA1 and BRCA2 variant(s) identified, are provided. If multiple variants are detected, the overall test interpretation most relevant to patient management is based on the most severe variant identified. Standard interpretative information included in test reports is listed below. Note that variants determined to have a classification of favor polymorphism or polymorphism are not included on the test report.

MyChoice CDx Components:

The overall results are composed of two major components, namely GIS Status, and tBRCA1/2 Status. The combined results form the basis of an overall interpretation of the MyChoice CDx Myriad HRD Status. Potential results for these two components are described below:

“GIS Status: Positive”
The test results demonstrate homologous recombination deficiency based on the GIS alone.

“GIS Status: Negative”
The test results demonstrate homologous recombination proficiency based on the GIS alone.

“Tumor Mutation BRCA1/BRCA2 Status (tBRCA1/2 Status): Positive for a Clinically Significant Mutation”
The test results demonstrate the presence of a deleterious or suspected deleterious sequencing mutation or large rearrangement.

“Tumor Mutation BRCA1/BRCA2 Status (tBRCA1/2 Status): Negative for a Clinically Significant Mutation”
The test results do not demonstrate the presence of a deleterious or suspected deleterious sequencing mutation or large rearrangement in BRCA1 or BRCA2. This category includes genetic variants for which published data demonstrate absence of substantial clinical significance and truncating mutations in BRCA2 that occur at and distal to amino acid 3.326 (Mazoyer S et al., Nature Genetics 1996, 14:253-254). It also includes variants in the protein-coding region that neither alter the amino acid sequence nor are predicted to significantly affect exon splicing, and base pair alterations in non-coding portions of the gene that have been demonstrated to have no pathogenic effect on the length or stability of the mRNA transcript. There may be uncommon genetic abnormalities that will not be detected by MyChoice CDx testing (see Limitations).
Interpretive Criteria

A positive Myriad HRD status is defined as the presence of deleterious or suspected deleterious mutation(s) in the BRCA1 and/or BRCA2 genes and/or a positive Genomic Instability Score. The interpretive criteria are based on the following classification categories:

“Deleterious”: Includes most nonsense and frameshift mutations that occur at/or before the last known deleterious amino acid position of the affected gene. In addition, specific missense mutations and non-coding intervening sequence (IVS) mutations are recognized as deleterious on the basis of data derived from linkage analysis of high-risk families, functional assays, biochemical evidence, statistical evidence, and/or demonstration of abnormal mRNA transcript processing. Deletions and duplications of an entire exon(s) identified by the Myriad CDx test may also be interpreted to be deleterious. Deleterious large genomic rearrangements include single exon and multi-exonic deletions that are out-of-frame. Out-of-frame single or multi-exonic duplications are classified as deleterious if the orientation is determined to be in tandem and head-to-tail. In-frame deletions/duplications are interpreted on an individual basis and the specific evidence supporting the classification of these mutations is included in the individual patient report.

“Suspected Deleterious”: Includes genetic variants for which the available evidence indicates a high likelihood, but not definitive proof, that the mutation is deleterious. The specific evidence supporting an interpretation will be summarized for individual variants in the patient report.

“Genetic Variant of Uncertain Significance”: Includes missense variants and variants that occur in analyzed intronic regions whose functional significance has not yet been determined, as well as nonsense and frameshift mutations that occur distal to the last known deleterious amino acid positions of the affected genes.

“Favor Polymorphism” and “Polymorphism”: Includes genetic variants for which available evidence indicates that the variant is highly unlikely to alter protein production and/or function. Variants of this type are not reported.

Change of interpretation and issuance of amended reports

The classification and interpretation of all variants identified in the assay reflect the current state of scientific understanding at the time the report is issued. In some instances, the classification and interpretation of such variants may change as new scientific information becomes available.

Whenever there is a change in the classification of a variant that affects the overall interpretation of a patient’s test result, an amended report will be provided by Myriad.
PERFORMANCE CHARACTERISTICS

Summary of Non-clinical Laboratory Studies

The specific performance characteristics of the MyChoice CDx assay were determined by studies using FFPE tumor samples. Samples were selected to evaluate a range of representative tumor BRCA1 and BRCA2 sequence variants (e.g., single nucleotide variants, insertions or deletions, and variants in homopolymers) and LRs (e.g., deletions and duplications affecting single and multiple exons) detected by the MyChoice CDx assay, as well as a representative range of GIS, as reflected in the device labeling.

1. Correlation with Orthogonal Reference Method (Accuracy)

The accuracy of the Myriad HRD Status determined by the MyChoice CDx Test was demonstrated using a validated Next Generation Sequencing (NGS)-based assay with a combination of non-clinical samples and FFPE clinical specimens from cancer patients enrolled in clinical trials from whom sufficient quantity and quality of DNA was available for testing with the NGS comparator assay. A total of 209 FFPE tumor specimen-derived DNA samples were tested with both the assays. Samples representing the following subgroups were tested in the study: 5 tBRCA1/2 Status positive / GIS Status negative, 71 tBRCA1/2 Status negative / GIS Status negative, 66 tBRCA1/2 Status positive / GIS Status positive, 61 tBRCA1/2 Status negative / GIS Status positive.

a. tBRCA1/2 Sequence Variant and LR Analytical Calls

A total of 1,733/1,733 valid BRCA1/2 sequence variant test calls were observed compared to the valid reference (comparator) BRCA1/2 sequence variant calls with ≥ 10% allele frequencies across all samples evaluated. This corresponds to a positive percent agreement (PPA) of 100%. Including variants with < 10% allele frequency, a total of 1,733/1,734 valid BRCA1/2 sequence variant calls were observed, corresponding to a PPA of 99.94% with a 95% lower confidence limit of 99.7267%. In addition, a total of 3,605,951/3,605,951 valid BRCA1/2 sequence non-variant base calls were observed, corresponding to a negative percent agreement (NPA) of 100% with a 95% lower confidence limit of 99.9999%. A total of 402/402 concordant valid BRCA1/2 LR calls were observed compared to the valid reference (comparator) BRCA1/2 LR calls across all samples evaluated. This corresponds to an overall percent agreement (OPA) of 100% for LR calls.

The results of the accuracy study were evaluated for three patient outcomes: (i) the tBRCA1/2 Status based on BRCA1 and BRCA2 sequence and LR analyses; (ii) the GIS Status based on the GIS; and (iii) the overall Myriad HRD Status based on the combined results of the tBRCA1/2 Status and GIS Status. The agreement between the MyChoice CDx device and the comparator (reference) assay is summarized below.

b. tBRCA1/2 Status Results

Concordance analysis of all 200 valid patient results produced by both the MyChoice CDx and Comparator assays resulted in an OPA of 100%.

c. GIS Status Results
Concordance analysis of all 206 valid patient results produced by both the MyChoice CDx and Comparator assays revealed a PPA of 98.5%, a NPA of 97.4%, and an OPA of 98.1%.

d. Myriad HRD Status Results

Concordance analysis of all 206 valid patient results from both the MyChoice CDx and Comparator assay revealed a PPA of 98.5%, a NPA of 98.6%, and an OPA of 98.5%.

2. Analytical Sensitivity

a. Limit of Blank (LoB)

Twenty-six (26) FFPE normal tissue samples were tested wherein all low frequency variants were expected to be spurious technical artifacts rather than true biological events. The distribution of allele frequencies was plotted. The frequencies of spurious variants were typically very low with the distribution decreasing very rapidly from 1% to 5%. There were no spurious variants with frequency above 5%.

One hundred thirty-six (136) FFPE tBRCA1/2 Status negative (wildtype) tissue samples were analyzed. These samples produced 2,357,861/2,357,861 (100%) concordant non-variant base calls, resulting in a false positive rate of 0% with a 95% upper confidence limit of 0.00013%. Of the 136 samples, 70 samples were both tBRCA1/2 Status negative / GIS Status negative. These 70 samples produced 1,213,621/1,213,621 (100%) concordant non-variant base calls, resulting in a false positive rate of 0% with a 95% upper confidence limit of 0.00025%. This retrospective analysis provides empirical data for setting a minimum allele frequency threshold of 5% to differentiate spurious background noise from real variants.

b. Limit of Detection (LoD)

i. tBRCA1/2 Sequence Variants

DNAs from four FFPE samples with known BRCA1/2 pathogenic sequence variants [BRCA1 c.181T>G (p.Cys61Gly), a single nucleotide variant, BRCA1 c.1961del, a < 10 bp deletion in an 8 bp homopolymer sequence, BRCA1 c.5266dupC, a < 10 bp insertion and BRCA2 c.9117_9117+11del, a ≥ 10 bp deletion] and FFPE tumor DNA samples without these BRCA1/2 variants (WT) were used to create simulated tumor:normal DNA samples with 10%, 8%, 6.5%, 5% and 2% allele frequencies of these mutations. Twenty replicates of each of the various DNA mixes were run and BRCA1/2 sequencing analytical calls were analyzed using CLSI’s EP17-A2 probit analysis methods to determine the LoD of these mutations.

The results of this study show that the LoDs of the four pathogenic sequence variants have different ranges of allele frequencies. The LoD of a single bp substitution was at 7.23% allele frequency. The LoD of a < 10 bp deletion in an 8 bp homopolymer sequence was 6.66%. The LoD for a < 10 bp insertion was 6.36%, and the LoD for a ≥ 10 bp deletion was 5.98%.

ii. tBRCA1/2 Large Rearrangements

Two FFPE tumor DNA samples, each carrying a different large rearrangement (LR): [BRCA1 del exon 8 LR exon and BRCA2 del exons 19–21 LR], were each mixed with
a FFPE DNA with no detectable BRCA1/2 LR (WT) to create tumor : normal samples with 50%, 40%, 30% and 10% allele frequencies of each LR. Ten replicates of each of the various DNA mixes were run and the concordance between the BRCA1/2 LR test calls and reference calls from each of the undiluted tumor samples was analyzed. The LoD for each LR was defined based on CLSI’s EP17-A2 guidance that recommends ≥95% concordant, positive LR calls.

The tBRCA1/2 LR portion of the MyChoice CDx assay’s LoD for the ≥ 3 exons LR is at 30% allele frequency, while the LoD for the 1-2 exons LR is at 50% allele frequency.

iii. GIS Status

Four FFPE matched tumor-normal samples were evaluated in this LoD study. DNAs extracted from each pair of FFPE matched tumor and normal samples were mixed to create five different tumor:normal DNA mixes at 40%, 30%, 20%, 10% and 0% tumor DNA content. Ten replicates of each of the tumor:normal mixes were run along with replicates of the undiluted tumor and normal DNAs. All the tumor:normal DNA mixes were tested using the MyChoice CDx assay and final GIS Status results were analyzed to assess the LoD of this assay.

The results from this LoD study show that the MyChoice CDx assay yields highly valid GIS results and GIS Status results for all samples at all tumor DNA content levels that produced valid results. Based on this study, the LoD of the GIS portion of the MyChoice CDx assay is at ~ 30% tumor DNA content.

3. Analytical Specificity

a. Interference (tBRCA1/2 Sequence Variant and LR Analytical Calls)

To evaluate the potential impact of three classes of substances (endogenous [hemoglobin, triglycerides], exogenous [tissue marking dye, paraffin wax], and method-specific interferents [ethanol, NaOH]) that can potentially interfere with the assay, this study evaluated seven FFPE specimens (5 tBRCA1/2 Status positive / GIS Status positive and 2 tBRCA1/2 Status negative / GIS status negative) representing single nucleotide variants, insertions or deletions of < 10 bp in length, insertions or deletions of ≥ 10 bp in length, homopolymer variants (5+ bp), and large rearrangements affecting ≥ 3 exons. The effects of potential interfering substances were tested at one or two replicates to determine if they would impact the MyChoice CDx device, and the results were compared to the control (no additional interferents) condition.

All treated samples across all six substances at the high test levels passed the acceptance criteria, with the exception of method-specific NaOH at the high test level (1.0 N), which failed as 10/14 (71.4%) tests successfully generated valid BRCA1/2 sequencing and LR results. However, 14/14 (100%) of the tests at the low 0.4 N NaOH level passed. All samples run under each condition produced valid positive or negative patient calls. The positive and negative patient calls were 100% concordant when compared to samples without additional interferents.

b. Interference (GIS)
All treated samples produced valid GIS for each potential interfering substance tested. In addition, all samples run under each condition produced valid positive or negative patient calls showing 100% concordance when compared to samples without additional interferents. Taken together, these results demonstrate that the tBRCA1/2 sequencing and GIS portions of the assay are minimally impacted or not impacted by the presence of any of the substances tested in this study.

Necrosis of ≥ 10% of the tumor area was observed in 11% (n = 66) of tumor samples from the QUADRA study. Only 3% of samples had necrosis in > 10% of the tumor area, and no samples were identified with necrosis involving > 60% of the tumor area. The standard procedure in Myriad’s Anatomic Pathology laboratory is to macro-dissect the fixed tumor tissues on slides to maximize tumor content, i.e., minimize the inclusion of non-tumor content including necrotic tissue. Thus, based on the above analysis, it was concluded that necrosis of ≤ 60% does not impact the MyChoice CDx assay results.

c. Carryover

The purpose of this study was to demonstrate that the MyChoice CDx test minimizes carryover across samples. Fourteen FFPE specimens (11 tBRCA1/2 Status positive / GIS Status positive, 1 tBRCA1/2 Status positive / GIS Status positive and 2 tBRCA1/2 Status negative / GIS Status negative) were processed consecutively through DNA extraction. The DNA from these 14 FFPE samples were then set up in a checkerboard pattern, alternating between low (50 ng) and high (200 ng) inputs. Two consecutive batches were set up in this pattern, with one checkboard pattern inverted, to assess intra-run (1st batch) and inter-run (2nd batch) carryover. An additional reference batch was run with all samples at 200 ng to compare for concordance. For both the intra-run and inter-run batches, all 14/14 samples produced complete, valid analytical calls that were 100% concordant for BRCA1 and BRCA2 sequence and LR calls, and all GIS were valid. In addition, the patient results of all samples run in both batches were 100% concordant.

Additionally, using the SNVs analyzed in the assay, carryover was quantified within and between batches. The highest intra-run carryover observed was 0.2% and the highest inter-run carryover observed was 0.1%. The average intra-run carryover observed was 0.14% and average inter-run carryover was 0.10%. The overall analytical results show that the MyChoice CDx device has very low intra-run and inter-run sample carryover and poses minimal risk to patient results.

d. Cross-Reactivity

Cross-reactivity studies for sequence-based assays are intended to differentiate between target analyte sequences and sequences generated from other sources. Three types of spurious sequences that could potentially be mistaken for target sequences, e.g., pseudogenes or other genomic regions that are highly homologous to targeted genes and regions, off-target regions that hybridize to hybridization baits or DNA sequences that carry-through the process, and process artifacts and low quality sequences. The purpose of this study is to demonstrate that captured sequences not originating from target regions do not materially affect the MyChoice CDx test.

Sequencing data for 7 FFPE tissue samples (4 tBRCA1/2 Status positive /GIS Status
positive, 1 tBRCA1/2 Status negative / GIS Status negative, 1 tBRCA1/2 Status negative / GIS Status positive, and 1 tBRCA1/2 Status unknown / GIS Status positive) were processed through MyChoice CDx in quadruplicate, yielding 28 total test results that were mapped to the human genome sequence to reveal off-target sequences due to pseudogene and cross-reactivity of hybridization baits (capture probes).

The MyChoice CDx test has two methods for mitigating the impact of pseudogenes. The capture baits were designed to minimize the capture of pseudogene regions, and the analysis algorithm identifies and excludes pseudogene-derived sequences as part of BRCA1/2 variant detection and LR calling.

No off-target sequences were found to affect the tumor BRCA1 and BRCA2 component of the device. Only 1.2% of SNVs used to calculate the GIS were affected by off-target sequences with a minimal effect on the reported score. This retrospective analysis suggests that cross-reactivity poses a minimal risk to the MyChoice CDx device.

4. Repeatability and Reproducibility

The repeatability and reproducibility of the MyChoice CDx was investigated by testing DNA extracted from FFPE clinical specimens. The purpose of these studies was to demonstrate that the MyChoice CDx assay generates highly reproducible BRCA1 and BRCA2 sequencing and LR analytical calls and GIS on the tested samples from five different studies over five periods of time. In the first three studies (Studies 1–3), a total of 18 unique samples were tested. All but two of these samples were tested at the DNA input amount of 200 ng per assay. The fourth study (Study 4) evaluated 7 FFPE tumor specimens from the first three studies at 50 ng DNA input. The samples were run in duplicate per run, over 6 runs, using 3 lots of reagents, 3 different sets of instruments, 6 different operators, and 3 different data reviewers. An additional study (Study 5) was performed using 5 unique FFPE samples with 18 replicates per sample divided across 9 independent runs over multiple days using 3 different sequencers and 3 lots of critical reagents. Thus, a total of 23 FFPE samples were evaluated (10 tBRCA1/2 Status positive / GIS Status positive, 2 tBRCA1/2 Status positive / GIS Status negative, 6 tBRCA1/2 Status negative / GIS Status positive, 3 tBRCA1/2 Status negative / GIS Status negative, 2 tBRCA1/2 Status unknown / GIS Status positive). The samples contained single nucleotide variants, insertions or deletions of < 10 bp in length, insertions or deletions of ≥ 10 bp in length, homopolymer variants (5+ bp), large rearrangements affecting 1-2 exons, and large rearrangements affecting ≥ 3 exons. The tested samples had a wide GIS range.

a. tBRCA1/2 Sequence Variant and LR Analytical Calls

All 228/228 samples and replicates tested from Studies 1–3 produced complete, valid BRCA1 and BRCA2 sequencing and LR analytical calls. Overall, 2,220/2,220 BRCA1 and BRCA2 sequence variant calls, 3,951,603/3,951,603 non-variant bases and 456/456 LR calls across all samples/replicates tested were 100% concordant. Study 4 analyzed 7 FFPE tumor samples at 50 ng DNA inputs, which had previously been evaluated at 200 ng DNA input levels in Studies 1-3. All samples and replicates run generated complete (100%) valid BRCA1 and BRCA2 sequence and LR calls. Altogether, all 708/708 BRCA1 and BRCA2 sequence variant calls, 1,450,393/1,450,393 non-variant bases and 168/168 LR calls across all samples/replicates tested at the 50 ng DNA input.
level were 100% concordant. All BRCA1 and BRCA2 sequence variant and LR calls from the 7 samples run at 50 ng DNA inputs were 100% concordant with analytical calls from the same samples run at 200 ng DNA inputs.

Study 5 tested five new additional ovarian tumor specimens that were tBRCA1/2 Status negative / GIS Status positive. The specimens were tested at the lowest DNA input of 30 ng with GIS Status being low positives. The MyChoice CDx device produced 700/702 concordant BRCA1/2 sequence variant calls, at mean allele frequency (MAF) ≥ 10%, resulting in a 99.7% PPA. There were two false negative calls in one sample in a single nucleotide variant with a MAF of 11.2%. When all valid BRCA1/2 sequence variants were analyzed, including those below 10% MAF, the MyChoice CDx device produced 788/792 concordant sequence variant calls resulting in a 99.5% PPA and 99.99% (1,553,331/1,553,332) concordance for non-variant base calls. For these five samples, BRCA1/2 LR concordance was 100% (165/165).

b. tBRCA1/2 Status Results

All samples and replicates run for all five studies produced valid positive or negative patient results, except for two samples, which were inconclusive for the tBRCA1/2 Status result. However, both of these samples had a positive GIS Status and were therefore both Myriad HRD Status positive. All valid positive and negative patient calls were 100% concordant, resulting in 100% PPA and NPA. In addition, valid patient results for all samples run at 50 ng DNA input level were 100% concordant with their corresponding sample run at the 200 ng DNA input level.

c. GIS

For the GIS portion of the assay, all 23 samples and replicates at all DNA inputs (200 ng, 50 ng, and 30 ng) from the five studies produced valid GIS. The overall 95% confidence interval for the true proportion of majority calls across samples correspond to 98.8% - 100% for the first four studies and 81.5% - 100% for the fifth study.

d. GIS Status and Myriad HRD Status Results

All samples and replicates run for all five studies produced complete, valid, positive, or negative patient calls in both sets of patient results. All positive and negative patient calls were 100% concordant, resulting in 100% PPA and NPA. In addition, all 7 samples run at the 50 ng and 200 ng DNA input levels were 100% concordant.

These results demonstrate that BRCA1/2 sequence variant and LR calls, GIS, and patient results of the MyChoice CDx assay are highly reproducible across the different process and reagent variabilities introduced in these studies.

5. Guardbanding

The MyChoice CDx guardband/robustness studies challenged the performance of the assay across three key parameters: (i) amount of FFPE tumor tissue-derived DNA input into the MyChoice CDx assay, (ii) hybridization temperature for probe capture, and (iii) library input onto the HiSeq instrument.

For the DNA input guardband study, 28 FFPE tumor samples (13 tBRCA1/2 Status positive / GIS Status positive, 1 tBRCA1/2 Status positive / GIS Status negative, 4 tBRCA1/2 Status
negative / GIS Status positive, 10 tBRCA1/2 Status negative / GIS Status negative) were run. For the hybridization temperature and library input guardband studies, seven FFPE tumor samples (3 tBRCA1/2 Status positive / GIS Status positive, 2 tBRCA1/2 Status positive / GIS Status negative, 2 tBRCA1/2 Status negative / GIS Status negative) were run in triplicate at standard MyChoice CDx assay conditions for the generation of reference results for each guardband condition. All analytical calls (i.e., BRCA1/2 sequence variants, LR calls, and GIS) and patient results for the reference and test samples and replicates were reported.

a. DNA Input Guardband/Robustness

Two studies were performed to evaluate the range of FFPE tumor extracted DNA input into the assay. In the first study, 14 samples were run in triplicate at 200 ng (used as the reference), 100 ng and 50 ng, and in singlet at 300 ng, 40 ng, 30 ng, 20 ng, and 10 ng. The second study ran a different set of 14 samples in triplicate at 200 ng (used as the reference) and 30 ng. The results from samples with different DNA input and samples near LoD of the tBRCA1/2 portion (< 10% and ≥ 10% MAF) were reported.

For the tBRCA1/2 portion of the MyChoice CDx assay, all valid BRCA1 and BRCA2 sequence variant calls from 300 ng to 10 ng DNA input levels were 99.8% concordant and all valid BRCA1 and BRCA2 non-variant base calls were 99.99% concordant. All valid BRCA1 and BRCA2 LR calls produced from 300 ng to 20 ng DNA input levels were 100% concordant (no valid LR calls were produced at the 10 ng input level). For the GIS portion of the MyChoice CDx assay, the total allowable error (TAE) analysis of valid GIS displayed acceptable amounts of bias and variation by passing pre-defined acceptance criteria from 300 ng to 30 ng input levels. In addition, the valid patient calls of all tests run were 100% concordant, except for a single false negative GIS Status patient call in one out of three replicates of one sample tested at 30 ng. All replicates of this sample had tBRCA1/2 Status positive patient results, and as such, all patient results were Myriad HRD Status positive.

b. Hybridization Temperature for Probe Capture

The hybridization temperature for probe capture was evaluated by varying the temperature by ± 1ºC and ± 2ºC from the 65ºC standard condition. Seven FFPE samples were run at standard hybridization temperature to generate reference results. The results of the 7 samples from each guardband test condition were compared to the results obtained at the standard hybridization temperature. All samples across all guardband test conditions generated complete, valid BRCA1 and BRCA2 sequencing and LR results, as well as valid GIS. All BRCA1 and BRCA2 sequence and LR calls were 100% concordant. Additionally, all samples tested at all guardband conditions produced valid GIS analytical results and all patient results were 100% concordant.

c. Library Input onto the HiSeq Instrument

The library input onto the HiSeq instrument was evaluated by varying the library concentration by ± 2 pM and ± 4 pM from the 6 pM standard condition. Seven FFPE samples were run at the standard library input amounts to generate reference results. The results of the 7 samples from each guardband test condition were compared to the results obtained at the standard library input amounts. Samples across all guardband test conditions generated complete, valid BRCA1 and BRCA2 sequencing and LR results, as
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well as GIS. All BRCA1 and BRCA2 sequence and LR calls were 100% concordant. All GIS generated at the different test conditions were valid and passed the pre-defined acceptance criteria. In addition, all patient results were 100% concordant.

These results show that the MyChoice CDx test is robust and is not affected by these process variations.

6. Stability Studies
   a. Stability of FFPE Clinical Specimens

   This study evaluated the real-time stability of FFPE tumor blocks and FFPE tumor sections on slides, stored at laboratory temperature. Reference results were defined from replicates of each specimen run at the initial (earliest) time point. The analytical and patient results of aged specimens, run in singlet, were compared to the reference calls for concordance.

   i. FFPE Tumor Blocks

   Stability data was analyzed for 16 unique FFPE tumor blocks for up to the 5.5-year time point. Analytical calls from aged blocks at each time-point were compared to those from the initial (T0), reference time point. Analytical results have been obtained for 12, 7 and 6 samples at the 3.5, 5.0, and 5.5 year time points, respectively. For BRCA1 and BRCA2 sequencing, a single false positive BRCA2 sequence variant was called at an allele frequency of 8.3% at the 3.5 year time point. All other BRCA1 and BRCA2 sequence variant calls were 100% concordant across all aged FFPE tumor blocks tested at each of the time points. All BRCA1 and BRCA2 LR calls were 100% concordant. In addition, the GIS were valid and patient results were 100% concordant across all aged blocks tested at the different time points. The stability study has been confirmed up to 5.5 years.

   ii. FFPE Tumor Sections

   The stability of unique FFPE tumor sections on slides is being evaluated at the following time points: 0, 1, 3, 5 and 5.5 years. Analytical calls from aged tumor sections at each testing time-point were compared to those from the initial (T0), reference time point. At the 1 year time point, all 10/10 samples generated complete, valid analytical calls and patient results. All BRCA1 and BRCA2 sequence and LR calls were 100% concordant. In addition, all GIS were valid and all patient results were 100% concordant across all specimens tested. The stability study has thus far been confirmed up to 1-year time point.

   b. FFPE Tumor Extracted DNA Stability

   The stability of DNAs extracted from 9 FFPE tumor specimens and stored at -20 ºC was evaluated at 0 (T0), 30 days, 60 days, 90 days and 6 months. Analytical calls from aged DNAs at each time-point were compared to those from the initial (T0), reference time point. All tests performed from 30 days to 6 months produced complete, valid analytical calls and patient results. For each of the stability testing time points, all BRCA1 and BRCA2 sequence and LR calls were 100% concordant. In addition, the GIS were valid and patient results were 100% concordant across all aged FFPE tumor extracted DNAs. The claimed stability for the extracted DNA is 5 months at -20°C.
c. Reagents Stability

The real-time stability of critical reagents used in the device was evaluated. Three lots of each of these reagents were stored at specified conditions and run at 0, 1 month, and 4 months. Each reagent lot was tested with 6 FFPE tumor DNAs in duplicate at each aged reagents time point. Analytical calls from aged reagents at each time-point were compared to those from the initial (T0) reference time point. All tests performed at 1 month and 4 months produced complete, valid analytical calls and patient results. For each of the stability testing time points, all BRCA1 and BRCA2 sequence and LR calls were 100% concordant, the GIS generated were valid, and all the patient results were 100% concordant across all lots of aged reagents.

7. Assay Migration

Myriad performed assay migration studies to support the migration of MyChoice CDx from the HiSeq 2500 sequencer (old system) to NovaSeq 6000 sequencer (new system). The studies included concordance between the new and old systems, precision, LoB, LoD and reagent stability with the new system.

Rates of test validity and analytical (e.g., BRCA1/2 sequencing, BRCA1/2 LR, and GIS) and patient status (i.e., tBRCA1/2 Status, GIS Status, and Myriad HRD Status) agreements were compared between the HiSeq 2500 and NovaSeq 6000 sequencers for the assay migration, precision, LoB, BRCA1/2 sequencing LoD, GIS Status LoD, and reagents stability studies for the subset of samples tested on both sequencers in each study. No statistically significant differences (α = 0.05, p ≤ 0.05) in the rates of test validity or analytical and patient status agreements were observed in any study, indicating equivalent MyChoice CDx performance using the HiSeq 2500 and NovaSeq 6000 sequencers.

The following sections report MyChoice CDx analytical performance using the NovaSeq 6000 sequencer.

a. Concordance between MyChoice CDx using NovaSeq 6000 Sequencer (new system) and MyChoice CDx using HiSeq 2500 Sequencer (old system)

The concordance of the Myriad HRD Status determined by the current MyChoice CDx Test using the NovaSeq 6000 sequencer (new system) compared to the MyChoice CDx Test using the HiSeq2500 sequencer (old system) was demonstrated using non-clinical fixed tumor samples from cancer patients. A total of 246 fixed tumor specimen-derived samples were tested with both the assays at four DNA inputs: 200 ng (185 samples), 100 ng (11 samples), 30 ng (25 samples), and 7.5 ng (25 samples). Samples representing the following subgroups were tested in the study: 5 tBRCA1/2 Status positive / GIS Status negative, 123 tBRCA1/2 Status negative / GIS Status negative, 62 tBRCA1/2 Status positive / GIS Status positive, 44 tBRCA1/2 Status negative / GIS Status positive.

i. tBRCA1/2 Sequence Variant and LR Analytical Calls

A total of 2,127/2,131 valid BRCA1/2 sequence variant Test calls were observed, corresponding to a PPA of 99.81% with a 95% confidence interval lower limit of 99.52%. In addition, a total of 4,247,898/4,247,907 valid BRCA1/2 sequence non-variant
base Test calls were observed, corresponding to a negative percent agreement (NPA) of 99.99979% with a 95% confidence interval lower limit of 99.99960%. A total of 440/440 concordant valid BRCA1/2 LR Test calls were observed, with 426/426 negative Test calls and 14/14 positive Test calls, compared to the old system BRCA1/2 LR calls across all samples evaluated. This corresponds to an overall percent agreement (OPA), NPA, and PPA of 100% for LR calls.

ii. **tBRCA1/2 Status Results**

Concordance analysis of all 243 valid patient results produced by both the new and old systems resulted in a NPA of 99.4%, PPA of 98.6%, and OPA of 99.2%.

iii. **GIS Status Results**

Concordance analysis of all 230 valid patient results produced by both the new and old systems resulted in a NPA of 98.4%, PPA of 97.1%, and an OPA of 97.8%.

iv. **Myriad HRD Status Results**

Concordance analysis of all 232 valid patient results from both the new and old system resulted in a NPA of 99.2%, a PPA of 96.4%, and an OPA of 97.8%.

b. **Analytical Sensitivity**

i. **LoB**

Fifty-three (53) FFPE normal, tBRCA1/2 Status negative tissue samples were tested in duplicate with the MyChoice CDx Test using the NovaSeq 6000 sequencer wherein all low frequency variants were expected to be spurious technical artifacts rather than true biological events. A false positive call rate of 0.00016%.

ii. **LoD**

1. **tBRCA1/2 Sequence Variants**

The sequence variant LoD claims using fixed tumor samples with representative pathogenic sequence variants (single base substitutions, insertions and/or deletions (indels), and homopolymers) was established for the MyChoice CDx Test using the NovaSeq 6000 sequencer at the lowest (7.5 ng) and highest (200 ng) DNA inputs. DNAs from fixed tumor samples containing the sequence variants were mixed with DNAs from unmatched (i.e., from different individuals) fixed normal (i.e., non-tumor) samples containing no sequence variants at the same gene location to generate contrived tumor:normal DNA mixes. These mixes were titrated at 4-5 different target variant allele frequencies and each tested with either 10 or 20 replicates. LoDs based on a 95% hit rate approach are presented for each sequence variant in Table 2.
Table 2: Summary of Sequence Variant LoDs

<table>
<thead>
<tr>
<th>Variant</th>
<th>Variant Type</th>
<th>LoD AF 200 ng DNA Input</th>
<th>LoD AF 7.5 ng DNA Input</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRCA1 c.3648del (p.Leu1216Phefs*19)</td>
<td>Indel &lt; 10 bp</td>
<td>5%</td>
<td>20%</td>
</tr>
<tr>
<td>BRCA1 c.5503C&gt;T (p.Arg1835*)</td>
<td>Single Base Substitution</td>
<td>8%</td>
<td>20%</td>
</tr>
<tr>
<td>BRCA1 c.4964_4982del (p.Ser1655Tyrfs*16)</td>
<td>Indel ≥ 10 bp</td>
<td>5%</td>
<td>15%</td>
</tr>
<tr>
<td>BRCA2 c.5350_5351del (p.Asn1784Hisfs*2)</td>
<td>Homopolymer +5 bp and Indel &lt; 10 bp</td>
<td>5%</td>
<td>10%</td>
</tr>
</tbody>
</table>

AF = allele frequency; Indel = insertion and/or deletion; bp = base pairs

2. tBRCA1/2 Large Rearrangements

The LR LoD claims using fixed tumor samples with representative LRs was established at the lowest (7.5 ng) and highest (200 ng) DNA inputs. DNAs from fixed tumor samples containing the LRs were mixed with DNAs from unmatched (i.e., from different individuals) fixed normal (i.e., non-tumor) samples containing no LRs at the same gene location to generate contrived tumor:normal DNA mixes. These mixes were titrated at 5 different target LR allele frequencies and each tested with 10 replicates. LoDs based on a 95% hit rate approach are presented for each LR in Table 3. A LoD could not be determined for the BRCA2 uncharacterized large rearrangement (ULR) at the 7.5 ng DNA input.

Table 3: Summary of Large Rearrangement LoDs

<table>
<thead>
<tr>
<th>LR</th>
<th>LR Type</th>
<th>LoD AF 200 ng DNA Input</th>
<th>LoD AF 7.5 ng DNA Input</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRCA1 del exons 9-10</td>
<td>1-2 exon LR</td>
<td>30%</td>
<td>50%</td>
</tr>
<tr>
<td>BRCA2 ULR</td>
<td>≥ 3 exon LR</td>
<td>30%</td>
<td>ND</td>
</tr>
</tbody>
</table>

AF = allele frequency; ND = not determined; ULR = uncharacterized large rearrangement

3. GIS Status

Two fixed tumor contrived tumor-normal samples, one GIS Status negative and one GIS Status positive, were evaluated in this LoD study for the MyChoice CDx Test using the NovaSeq 6000 sequencer. DNAs extracted from each pair of matched (i.e., from the same individual) fixed tumor and normal (i.e., non-tumor) samples were mixed to create at five tumor:normal DNA mixes ranging from 49% to 10% tumor DNA content and run at both 200 ng and 7.5 ng DNA input. Ten to 20 replicates of each of the tumor:normal mixes were run along with replicates of the undiluted tumor and normal DNAs. All the tumor:normal DNA mixes were tested using the
MyChoice CDx Technical Information

MyChoice CDx assay and GIS Status results were analyzed to assess the LoD of this assay.

LoDs based on a 95% hit rate approach are presented for each GIS Status in Table 4. The LoD of the GIS Status negative sample was ~30% tumor content at 200 ng DNA input and could not be determined at 7.5 ng DNA input. The LoD of the GIS Status positive sample was ~30% tumor content at 200 ng DNA input and ~49% tumor content at 7.5 ng DNA input.

<table>
<thead>
<tr>
<th>GIS Status</th>
<th>LoD Tumor Content 200 ng DNA Input</th>
<th>LoD Tumor Content 7.5 ng DNA Input</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>30%</td>
<td>ND</td>
</tr>
<tr>
<td>Positive</td>
<td>30%</td>
<td>49%</td>
</tr>
</tbody>
</table>

ND = not determined

c. **Precision**

i. **Repeatability and Reproducibility**

A total of 14 samples were tested in 18 replicates each using the MyChoice CDx Test using the NovaSeq 6000 sequencer. The samples were run in duplicate per run, using 3 lots of reagents, 3 different sets of instruments, 3 sets of operators, and 3 sets of data reviewers, at both 200 ng and 7.5 ng DNA input, for a total of 18 replicates per sample per DNA input. In total, 5 tBRCA1/2 Status positive / GIS Status positive, 2 tBRCA1/2 Status positive / GIS Status negative, 3 tBRCA1/2 Status negative / GIS Status positive, 4 tBRCA1/2 Status negative / GIS Status negative samples were tested. The samples contained 127 single nucleotide variants, nine insertions or deletions of < 10 bp in length, two insertions or deletions of ≥ 10 bp in length, eight homopolymer variants (5+ bp), two large rearrangements affecting 1-2 exons, two large rearrangements affecting ≥ 3 exons, and a range of GISs.

1. **tBRCA1/2 Sequence Variant and LR Analytical Calls**

252 samples/replicates tested at the 200 ng DNA input produced valid BRCA1 and BRCA2 sequencing and LR analytical results. 250 samples/replicates tested at the 7.5 ng DNA input produced valid BRCA1 and BRCA2 sequencing and LR analytical results. When the agreement of valid BRCA1/2 sequence variant analytical calls produced for variants with a reference mean allele frequency (MAF) ≥ 10% was assessed, the samples evaluated at the 200 ng DNA input had a PPA of 2,358/2,358 (100%), while the samples evaluated at the 7.5 ng DNA input had a PPA of 2,338/2,343 (99.79%). PPA for all valid sequence variant calls (including those with a reference MAF < 10%) were 2,484/2,484 (100%) at 200 ng DNA input and 2,440/2,469 (98.83%) at 7.5 ng DNA input.
NPA for valid BRCA1/2 sequence non-variant base calls were 4,404,685/4,404,690 (99.99989%) at 200 ng DNA input and 4,369,692/4,369,727 (99.99920%) at 7.5 ng DNA input. For LRs, at the 200 ng DNA input, MyChoice produced 72/72 (100%) PPA, 426/426 (100%) NPA, and 498/498 (100%) OPA for valid BRCA1/2 LR calls. At the 7.5 ng DNA input, MyChoice produced 70/70 (100%) PPA, 422/422 (100%) NPA, and 492/492 (100%) OPA for valid BRCA1/2 LR calls.

2. tBRCA1/2 Status Results

252 samples/replicates tested at the 200 ng DNA input produced valid positive or negative tBRCA1/2 Status patient results. 250 samples/replicates tested at the 7.5 ng DNA input produced valid positive or negative tBRCA1/2 Status patient results. MyChoice produced a 126/126 (100%) PPA, 126/126 (100%) NPA, and 252/252 (100%) OPA at the 200 ng DNA input. MyChoice produced a 125/125 (100%) PPA, 124/125 (99.2%) NPA, and 249/250 (99.6%) OPA for the tBRCA1/2 Status patient results at the 7.5 ng DNA input.

3. GIS

For the GIS portion of the assay, 252 samples/replicates produced valid GIS at the 200 ng DNA input, and 229 samples/replicates produced valid GIS at the 7.5 ng DNA input.

4. GIS Status and Myriad HRD Status Results

For the GIS Status patient results, MyChoice produced a 144/144 (100%) PPA, 108/108 (100%) NPA, and 252/252 (100%) OPA at the 200 ng DNA input. At the 7.5 ng DNA input, MyChoice produced a 139/139 (100%) PPA, 90/90 (100%), NPA, and 229/229 (100%) OPA.

For the overall Myriad HRD Status patient results, MyChoice produced 180/180 (100%) PPA, 72/72 (100%) NPA, and 252/252 (100%) OPA at the 200 ng DNA input, and 178/178 (100%) PPA, 62/63 (98.4%) NPA, and 240/241 (99.6%) OPA at the 7.5 ng DNA input.

d. Stability Studies

i. Reagents Stability

Multiple real-time studies have been performed to assess the stability of critical reagents that have been updated over time. Two studies support the stability of current critical reagents.

In the first study, which used the HiSeq 2500 (old system), three lots of all critical reagents, including an updated capture probes library, were stored at specified conditions and run with 6 fixed tumor DNAs in duplicate at T0, and at 2-month, 9-month, and 15-month stability time points. For each of the stability time points, all BRCA1 and BRCA2 sequence and LR calls were 100% concordant, the GIS generated were valid, and all the tBRCA1/2 Status, GIS Status, and Myriad HRD Status patient results had 100% OPA.
across all lots of aged reagents.

In the second study, which used the NovaSeq 6000 (new system), three lots of critical sequencing reagents were stored at specified conditions and run with 6 fixed tumor derived samples in duplicate at T0 and at a 2-month stability time point. At the 2-month stability time point, all BRCA1 and BRCA2 sequence and LR calls were 100% concordant, the GIS generated were valid, and all the tBRCA1/2 Status, GIS Status, and Myriad HRD Status patient results had 100% OPA across all lots of aged reagents. This stability study is ongoing.

Summary of Clinical Studies


   The niraparib clinical study PR-30-5017-C (PRIMA) was a randomized, double-blind, placebo-controlled, multicenter Phase 3 study design in subjects with ovarian, fallopian tube, and primary peritoneal cancer, collectively referred to as ovarian cancer. The objective of PRIMA was to evaluate the therapeutic effect of maintenance niraparib treatment following response to first-line platinum-based chemotherapy in patients with advanced ovarian cancer.

   a. Accountability of PMA Cohort

      A total of 733 patients were screened into the PRIMA study for homologous recombination deficiency (HRD) testing. Test results were required prior to randomization. Subjects with a documented deleterious germline (gBRCA) or somatic (sBRCA) BRCA mutation by local results were considered to have homologous recombination deficient tumors for stratification and randomization purposes; the tumor test was performed concurrently to confirm local results. Of these 733 randomized patients, 487 niraparib-treated and 246 placebo-treated patients were included in the PMA cohort. The retrospective analysis included the testing of 713 ovarian FFPE tumor specimens (7 samples were retests from existing patients and 4 samples from 4 patients who were identified as screen failures after re-matching ID with clinical database) from PRIMA, where biomarker calls from the CTA were compared with those from the CDx. Therefore, the final PMA cohort by CTA/CDx includes a total of 733 patients: 373 HRD positive patients, 249 HRD negative patients, 80 patients who had inconclusive results from HRD testing, and 31 patients without sufficient sample for HRD testing.

   b. Effectiveness Results

      The primary endpoint was PFS, defined as the time from treatment randomization to the earlier date of assessment of progression (by BICR) or death by any cause in the absence of progression. PFS was based on radiology assessment using RECIST v1.1 criteria. Efficacy analyses were primarily conducted on the intent to treat (ITT) population, in the overall and homologous recombination deficient populations.

      The main efficacy conclusions for the prespecified primary analysis population were as follows:
The study met its primary efficacy objective; treatment with niraparib prolonged median PFS by 11.5 months compared to placebo in subjects with homologous recombination deficient advanced ovarian cancer following response to front-line platinum-based chemotherapy. Median PFS as determined by BICR based on RECIST (version 1.1) was 21.9 months in the niraparib arm and 10.4 months in the placebo arm (HR 0.43 [95% CI: 0.310, 0.588]; p<0.0001).

In the overall population, treatment with niraparib prolonged median PFS by 5.6 months compared to placebo. Median PFS as determined by BICR based on RECIST (version 1.1) was 13.8 months in the niraparib arm and 8.2 months in the placebo arm (HR 0.62 [95% CI: 0.502, 0.755]; p<0.0001).

### Table 5: Efficacy Results*

<table>
<thead>
<tr>
<th></th>
<th>HRD Positive Population</th>
<th>Overall Population</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ZEJULA (N=247)</td>
<td>Placebo (N=126)</td>
</tr>
<tr>
<td></td>
<td>Placebo (N=487)</td>
<td>Placebo (N=246)</td>
</tr>
<tr>
<td>PFS events, n (%)</td>
<td>81 (33)</td>
<td>73 (58)</td>
</tr>
<tr>
<td></td>
<td>232 (48)</td>
<td>155 (63)</td>
</tr>
<tr>
<td>PFS Median (95% CI),</td>
<td>21.9 (19.3, NE)</td>
<td>10.4 (8.1, 12.1)</td>
</tr>
<tr>
<td>in months</td>
<td>13.8 (11.5, 14.9)</td>
<td>8.2 (7.3, 8.5)</td>
</tr>
<tr>
<td>Hazard Ratio (HR)c</td>
<td>0.43 (0.31, 0.59)</td>
<td>0.62 (0.50, 0.76)</td>
</tr>
<tr>
<td>(95% CI)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>p-valued</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

*a efficacy analysis was based on blinded independent central review (BICR).
*b In the HR proficient (HRD negative) population (N=249), a hazard ratio of 0.68 (95% CI [0.49, 0.94]) was observed.
*In the HR not determined (HRnd) population (N=111), a hazard ratio of 0.85 (95% CI [0.51, 1.43]) was observed.
*c based on a stratified Cox proportional hazards model
*d based on a stratified log-rank test

2. Summary of Clinical Study - Olaparib NCT03737643 (PAOLA-1) for maintenance treatment.

The olaparib clinical study was a randomized, double-blind, placebo-controlled, multi-center trial that compared the efficacy of Lynparza in combination with bevacizumab versus placebo/bevacizumab for the maintenance treatment of advanced high-grade epithelial ovarian cancer, fallopian tube or primary peritoneal cancer following first-line platinum-based chemotherapy and bevacizumab.

a. Accountability of PMA Cohort

Eight hundred and six (806) patients were randomized (2:1) to receive Lynparza tablets 300 mg orally twice daily in combination with bevacizumab (n=537) or placebo/bevacizumab (n=269) who had no evidence of disease (NED) due to complete surgical resection, or who were in complete response (CR), or partial response (PR) following completion of first-line platinum-containing chemotherapy and bevacizumab. Randomization was stratified by first-line treatment outcome (timing and outcome of cytoreductive surgery and response to platinum-based chemotherapy) and tBRCAm status, determined by prospective local testing.
Available tumor samples were retrospectively tested using the MyChoice® CDx test to determine HRD positive status, defined by either a deleterious or suspected deleterious BRCA mutation, or a genomic instability score ≥42. 46.7% of patients (376/806) were HRD status positive (26.7% [215/806] were tBRCAm and 20.0% [161/806] were non-tBRCAm) and 33.1% (267/806) were HRD status negative.

A total of 20.2%, (163/806) of patients had an unknown Myriad HRD status (11.5% had no samples available for testing and 8.7% had a cancelled or failed test).

b. Effectiveness Results

The major efficacy outcome measure was investigator-assessed PFS evaluated according to RECIST, version 1.1. PAOLA-1 demonstrated a statistically significant improvement in investigator-assessed PFS for Lynparza/bevacizumab compared to placebo/bevacizumab (HR 0.33; 95% CI 0.25-0.45 with a median of 37.2 months for Lynparza/bevacizumab vs 17.7 months for placebo/bevacizumab).

Efficacy results from a biomarker subgroup analysis of 387 patients with HRD positive tumors, identified post-randomization using the Myriad MyChoice® HRD Plus tumor test, who received Lynparza/bevacizumab (n=255) or placebo/bevacizumab (n=132), are summarized in Table 6. Results from a blinded independent review of PFS were consistent.

Table 6: Efficacy Results – PAOLA-1 (HRD positive status*, Investigator Assessment)

<table>
<thead>
<tr>
<th></th>
<th>Lynparza/bevacizumab (n=255)</th>
<th>Placebo/bevacizumab (n=132)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Progression-Free Survival</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of events (%)</td>
<td>87 (34%)</td>
<td>92 (70%)</td>
</tr>
<tr>
<td>Median, months</td>
<td>37.2</td>
<td>17.7</td>
</tr>
<tr>
<td>Hazard ratio* (95% CI)</td>
<td>0.33 (0.25, 0.45)</td>
<td></td>
</tr>
</tbody>
</table>

* Median follow-up of 24.4 months in both treatment arms for censored patients.

The analysis was performed using an unstratified Cox proportional hazards model.

CI Confidence interval

3. Summary of Clinical Study – Olaparib D0818C00001 (SOLO1)

The olaparib clinical study D0818C00001 (SOLO1) was a Phase III, randomised, double blind, placebo controlled, multicenter study to assess the efficacy of olaparib maintenance monotherapy in advanced ovarian cancer patients (including patients with primary peritoneal and/or fallopian tube cancer) who had responded following first-line platinum based chemotherapy and carrying BRCA mutations (documented mutation in BRCA1 or BRCA2) that were predicted to be deleterious or suspected deleterious (known or predicted to be detrimental/lead to loss of function). A total of 391 patients were enrolled from 118 centers in 15 countries, including the United States. Patients were required to have documented evidence of a deleterious or suspected deleterious mutation in either BRCA1 or BRCA2 to be
enrolled into the study. Evidence of a qualifying BRCA mutation could be from either an existing BRCA mutation result from local testing, or from prospective testing performed by Myriad (Salt Lake City, UT) using the Myriad Integrated BRACAnalysis® or for Chinese patients by BGI Clinical Laboratories (Shenzhen, China) using a local validated test.

A post-randomisation analysis of a subset of the SOLO1 tumor samples was performed to identify the population that would be defined as Tumor BRCA1/2 Mutation Status (tBRCA1/2) positive by the Myriad MyChoice® CDx test. The clinical performance of the MyChoice CDx test for the SOLO1 study was based on the Tumor Mutation BRCA1/BRCA2 (tBRCA1/2) Status and was established based on available residual extracted DNA generated from tumor FFPE samples collected from patients enrolled in the SOLO1 study. In the patient population, the Myriad BRACAnalysis CDx test detected 15 pathogenic large rearrangements (LRs). Large rearrangement analysis could not be reported by the Myriad MyChoice CDx test for two of the 15 pathogenic LRs due to poor quality of the tumor specimens. Of the remaining 13 LRs detected by the BRACAnalysis CDx test, 12 were also detected by the MyChoice CDx test. One pathogenic LR was not reported using the MyChoice CDx test due to a known limitation of the device. Three of the 12 LRs were detected by both BRACAnalysis CDx and MyChoice CDx. However, differences in interpretation criteria between the germline and tumor-based tests led to differences in the reported names and classifications of these LRs.

a. Accountability of PMA Cohort

Out of the 391 patients randomized in the SOLO1 study, extracted FFPE DNA samples from 333 patients were available for retrospective MyChoice® CDx testing. Samples from 298 patients were tested using MyChoice® CDx test, and valid tBRCA1/2 results were reported for 292 patients. Among those, 284 patients were confirmed to carry a deleterious or suspected deleterious tBRCA mutation and 8 patients were not confirmed to carry a deleterious or suspected deleterious tBRCA mutation by the MyChoice® CDx test. The PMA cohort represented 74.7% (292/391) of the full analysis set in SOLO1.

b. Effectiveness Results

The primary objective of this study was to determine the efficacy by PFS (using investigator assessment of scans according to modified Response Evaluation Criteria in Solid Tumors [RECIST] 1.1) of olaparib maintenance monotherapy compared with placebo in BRCA mutated high risk advanced ovarian cancer patients who are in clinical CR or PR following first line platinum based chemotherapy. The study population consisted of 391 ovarian cancer patients with a deleterious or suspected deleterious germline BRCA mutation as detected by the Myriad Integrated BRACAnalysis® test, BRACAnalysis CDx® test, BGI test or local test results. A statistically significant and clinically relevant improvement in investigator-assessed PFS, as evidenced by the magnitude of effect: a 70% reduction in the risk of disease progression or death at any point in time for olaparib vs placebo treated patients (HR 0.30; 95% CI 0.23-0.41; p<0.0001), with a median PFS not reached in the olaparib arm vs 13.8 months for placebo. Median follow-up for PFS defined as time from randomisation to date of censoring was approximately 41 months on both arms.
The effectiveness of the tBRCA1/2 Status of Myriad MyChoice® CDx test was based on a subset of 284 confirmed ovarian cancer patients with deleterious or suspected deleterious tBRCA1/2 mutations.

The clinical outcome data for the 284 patients with a confirmed tBRCA1/2 mutation was as follows: a 71% reduction in the risk of disease progression or death at any point in time for olaparib vs placebo treated patients (HR 0.29; 95% CI 0.21-0.41; p<0.0001), with a median PFS not reached in the olaparib arm vs 13.8 months for placebo. Median follow-up for PFS defined as time from randomisation to date of censoring was approximately 41 months on both arms. Taken together, these results are very similar to those observed in the 391 patients in the SOLO1 study, which supports the effectiveness of the device.

The clinical outcome results for cases classified as having a deleterious or suspected deleterious tumor BRCA mutation by the Myriad MyChoice® CDx test are shown in the table below.

<table>
<thead>
<tr>
<th></th>
<th>SOLO1</th>
<th>Myriad MyChoice® CDx test</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Olaparib 300 mg bd a</td>
<td>Placebo</td>
</tr>
<tr>
<td><strong>PFS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of events: total number of patients (%)</td>
<td>102/206 (39)</td>
<td>96/131 (73)</td>
</tr>
<tr>
<td>Median PFS (months)</td>
<td>not reached</td>
<td>13.8</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.30 (0.23-0.41)</td>
<td></td>
</tr>
<tr>
<td>P-value (2-sided)</td>
<td>&lt;0.0001</td>
<td></td>
</tr>
</tbody>
</table>

a- tablet formulation