MyChoice® CDx Plus

HRD Companion Diagnostic Test

Identify the right ovarian cancer patients for **PARPi treatment** in time



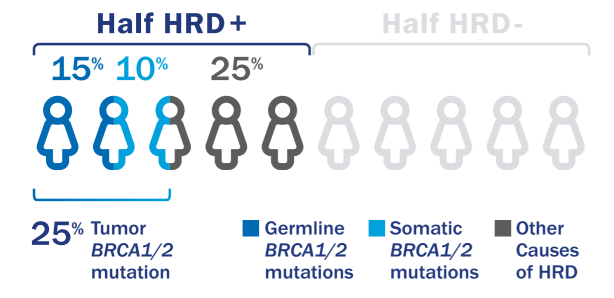
Missing out on a PARPi eligible woman

Optimizing HRD status evaluation

Finding women with HRD+ is complex

- In the early setting, ovarian cancer will progress in 7 out of 10 women.¹
- PARP inhibitors can slow down, potentially cure the disease, and help these women live longer. This works best for women with a certain type of ovarian cancer known as HRD+.^{2,3}
- One in four of these women have a change in their *BRCA1/2* genes. But 10% of these changes cannot be found by tests that look for inherited changes.⁴

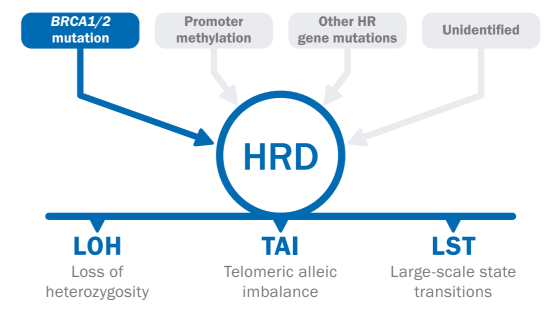
For about half of the women with HRD+ we are not able to determine their status by looking at each possible cause one by one.⁵



Combining multiple factors

- BRCA status and Genomic Instability Score combined identifies more women with a HRD+ tumor than any other test.
- BRCA1 and BRCA2 status is determined by changes in the sequence variants and identification of large rearrangements in the genes.
- The Genomic Instability Score is calculated based on the loss of heterozygosity, the telomeric imbalance, and large-scale transitions within the entire genome.

By combining these factors, you can find up to 3.5 times more patients with HRD+ than *BRCA* germline tests.⁶



Introducing MyChoice



Make sure that you don't miss a HRD+ patient

By combining tumor *BRCA* mutation testing and Genomic Instability Score, MyChoice identifies more patients eligible for PARP inhibitor treatment.



MyChoice provides a clear HRD status

Despite the very complex technology utilised, MyChoice gives you an easy to interpret result report for a clear HRD Status: positive or negative.



HRD Companion Diagnostic Test

Genomic Instability Score status

Tumor BRCA and Genomic Instability Score factors into final HRD status

Myriad Genetics HRD stat

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Genomic Instability Score status	Tumor BRCA status	Final Myriad Genetics HRD status	
Positive	Negative	Positive	
Positive	Positive Positive		
Negative	Positive	Positive	
Negative	Negative	Negative	

Identifying the tumor BRCA1 and BRCA2 status



MyChoice identifies and classifies BRCA1/2 sequence variants and large rearrangements as well as somatic and germline variants present in the tumor.

Assessing the Genomic Instability Score (GIS)



The GIS status is calculated with LOH (loss of heterozygosity), TAI (telomeric alleic imbalance), and LST (large-scale transitions) in the entire genome.



Get the results in 2-3 weeks

MyChoice delivers results in 2-3 weeks to aid you in making informed treatment decisions in a timely manner for your patients.

MyChoice HRD status includes BRCA & GIS



By looking at both the cause (BRCA) and the consequence (GIS) of the HRD status,
MyChoice is the most comprehensive test available.

First-in-class HRD test

Mentioned in major guidelines...



European Society of Medical Oncology

Validated scar-based HRD tests can be used to guide PARP inhibitor treatment. ESMO recognizes MyChoice CDx is the only scar based HRD test validated in the first-line maintenance setting.⁷



American Society of Clinical Oncology

Myriad MyChoice CDx is included in the new recommendations from The American Society of Clinical Oncology (ASCO) on the use of PARP inhibitors for the treatment and management of certain patients with advanced ovarian cancer.8



National Comprehensive Cancer Network®

Somatic testing should prioritize identification of molecular alterations that inform the use of effective interventions. This includes assessing BRCA1/2, loss of heterozygosity (LOH), or homologous recombination deficiency (HRD) status in the absence of a germline BRCA mutation.9

Superior to other solutions

		Many other HRD solutions	MyChoice
BRCA1 & BRCA2 status	Sequence variants	√	\checkmark
	Large rearrangement	0	✓
Prospectively validated genomic instability status	Loss of heterozygosity (LOH)	⊘ *	\checkmark
	Telomeric allelic imbalance (TAI)	0	\checkmark
	Large-scale state transition (LST)	0	✓

Not all tumor tests detect large rearrangements

MyChoice CDx conducts a comprehensive assessment of LOH, TAI, and LST across the entire genome

...with Level of Evidence 1A

PAOLA The Phase III PAOLA-1/ENGOT-ov25 trial evaluated maintenance olaparib plus bevacizumab in patients with newly diagnosed advanced ovarian cancer.²

Progression-free survival (PFS) in patients with HRD after 5 years median follow-up¹⁰



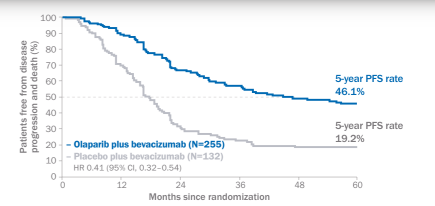
Reduction in risk of disease progression or death

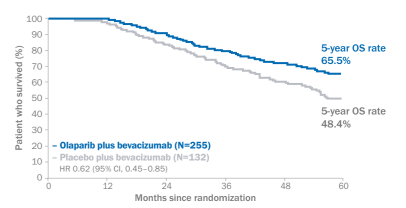
Final overall survival (OS) in patients with HRD after 5 years median follow-up¹⁰



Reduction in risk of death

for olaparib + bevacizumab vs bevacizumab alone



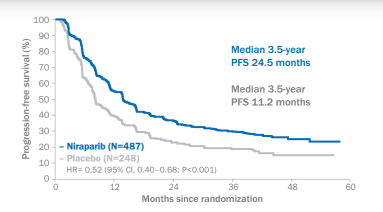


PRIMA The Phase III PRIMA study evaluated niraparib first-line maintenance therapy in patients with newly diagnosed advanced ovarian cancer after a response to first-line, platinum-based chemotherapy.³

Progression-free survival in patients with HRD with 3.5 years median follow-up¹¹



Reduction in risk of disease progression or death



 $[\]ensuremath{^{\star}}$ One commercially available test with prospectively validated %LOH

MyChoice®CDx Plus

HRD Companion Diagnostic Test

How to access MyChoice CDx Plus





MyChoice lab



Send MyChoice CDx Plus

request to your lab



Your lab send tissue to MyChoice lab



MyChoice lab analyses tissue



Result report is sent to you and your lab

Myriad Genetics is committed to illuminating the treatment pathway for every woman

Myriad Genetics is a leading genetic testing and precision medicine company dedicated to advancing health and wellbeing for all, empowering individuals with vital genetic insights and enabling doctors to better detect, treat and prevent disease.

Myriad discovers and provides genetic and genomic tests that determine the risk of developing disease, assess the risk of disease progression, and guide treatment decisions across medical specialties where critical genetic insights can significantly improve patient care and provide access to the right treatments.

References:

1. Ledermann, J. et al. Annals Of Oncology 2013 2. Ray-Coquard, I. et al. New England Journal Of Medicine 2019 3. Gonzalez-Martin, A. et al. New England Journal Of Medicine 2019 4. The Cancer Genome Atlas Research Network. Nature 2011 5. Watkins, J. A. et al. Breast Cancer Research 2014 6. Timms, K. M. et al. Journal of Clinical Oncology 2020 7. Miller R. E. et al. Annals of Oncology 2020 8. Tew W. P. et al. Journal of Clinical Oncology 2020 and 2022 9. Referenced with permission from the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) for Ovarian Cancer V.1.2023 10. Ray-Coquard I. et al. Annals of Oncology 2023 11. Gonzalez-Martin, A. et al. European Journal of Cancer 2023.





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The Myriad Genetics MyChoice® CDx Plus test was developed and performance characteristics were determined by Myriad Genetic Laboratories, Inc. and in compliance to In-Vitro Diagnostic Device Directive (98/79/EC) and is CE marked. Myriad is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-88) as qualified to perform high complexity clinical laboratory testing. Myriad is compliant with multiple international standards including, ISO 13485:2016 and ISO 15189: 2012 as applicable.

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