

TUMOR TYPE
Breast cancer (NOS)
COUNTRY CODE

REPORT DATE
21 May 2021
ORDERED TEST #
ORD-1095717-01

**ABOUT THE TEST** FoundationOne®Liquid CDx is a next generation sequencing (NGS) assay that identifies clinically relevant genomic alterations in circulating cell-free DNA

#### **PATIENT**

**DISEASE** Breast cancer (NOS)

DATE OF BIRTH 31 December 1971

SEX Female

MEDICAL RECORD # Not given

#### **PHYSICIAN**

MEDICAL FACILITY Arias Stella ADDITIONAL RECIPIENT None MEDICAL FACILITY ID 317319 PATHOLOGIST Not Provided

#### **SPECIMEN**

SPECIMEN ID 09656751 31-12-71 SPECIMEN TYPE Blood DATE OF COLLECTION 10 May 2021 SPECIMEN RECEIVED 14 May 2021

## Biomarker Findings

Blood Tumor Mutational Burden - 1 Muts/Mb Microsatellite status - MSI-High Not Detected Tumor Fraction - Cannot Be Determined

# **Genomic Findings**

For a complete list of the genes assayed, please refer to the Appendix.

**FGF6** R62H

0 Therapies with Clinical Benefit

O Clinical Trials

O Therapies with Lack of Response

### **BIOMARKER FINDINGS**

# Blood Tumor Mutational Burden - 1 Muts/Mb

Microsatellite status - MSI-High Not Detected

Tumor Fraction - Cannot Be Determined

#### THERAPY AND CLINICAL TRIAL IMPLICATIONS

No therapies or clinical trials. See Biomarker Findings section

MSI-High not detected. No evidence of microsatellite instability in this sample (see Appendix section).

Tumor fraction is an estimate of the percentage of circulating-tumor DNA (ctDNA) present in a cell-free DNA (cfDNA) sample based on observed aneuploid instability.

# No therapies or clinical trials are associated with the Genomic Findings for this sample.

If you have questions or comments about this result, please contact your Foundation Medicine customer support representative.

**Phone:** 1-888-988-3639

Online: foundationmedicine.com

Email: client.services@foundationmedicine.com

### GENOMIC FINDINGS WITH NO REPORTABLE THERAPEUTIC OR CLINICAL TRIALS OPTIONS

For more information regarding biological and clinical significance, including prognostic, diagnostic, germline, and potential chemosensitivity implications, see the Genomic Findings section.

FGF6 - R62H p. 5





TUMOR TYPE

Breast cancer (NOS)

COUNTRY CODE

DE

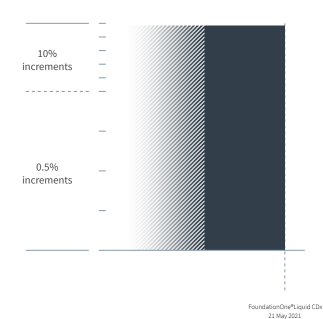
REPORT DATE
21 May 2021
ORDERED TEST #
ORD-1095717-01

NOTE Genomic alterations detected may be associated with activity of certain approved therapies; however, the therapies listed in this report may have varied clinical evidence in the patient's tumor type. Therapies and the clinical trials listed in this report may not be complete and/or exhaustive. Neither the therapies nor the trials identified are ranked in order of potential or predicted efficacy for this patient, nor are they ranked in order of level of evidence for this patient's tumor type. This report should be regarded and used as a supplementary source of information and not as the single basis for the making of a therapy decision. All treatment decisions remain the full and final responsibility of the treating physician and physicians should refer to approved prescribing information for all therapies. Therapies contained in this report may have been approved by the US FDA or other national authorities; however, they might not have been approved in your respective country. In the appropriate clinical context, germline testing of APC, BRCA1, BRCA2, BRIP1, MEN1, MLH1, MSH2, MSH6, MUTYH, NF2, PALB2, PMS2, PTEN, RAD51C, RAD51D, RB1, RET, SDHA, SDHB, SDHC, SDHD, SMAD4, STK11, TGFBR2, TP53, TSC1, TSC2, VHL, and WT1 is recommended.

Variant Allele Frequency is not applicable for copy number alterations.

FOUNDATIONONE®LIQUID CDx





| HISTORIC PATIENT FINDINGS        |        | ORD-1095717-01<br>VAF% |  |  |
|----------------------------------|--------|------------------------|--|--|
| Blood Tumor<br>Mutational Burden |        | 1 Muts/Mb              |  |  |
| Microsatellite status            |        | MSI-High Not Detected  |  |  |
| Tumor Fraction                   |        | Cannot Be Determined   |  |  |
| FGF6                             | ● R62H | 50.5%                  |  |  |

NOTE This comparison table refers only to genes and biomarkers assayed by prior FoundationOne®Liquid CDx, FoundationOne®Liquid, FoundationOne®, or FoundationOne®CDx tests. Up to five previous tests may be shown

For some genes in FoundationOne Liquid CDx, only select exons are assayed. Therefore, an alteration found by a previous test may not have been confirmed despite overlapping gene lists. Please refer to the Appendix for the complete list of genes and exons assayed. The gene and biomarker list will be updated periodically to reflect new knowledge about cancer biology.

As new scientific information becomes available, alterations that had previously been listed as Variants of Unknown Significance (VUS) may become reportable.

Tissue Tumor Mutational Burden (TMB) and blood TMB (bTMB) are estimated from the number of synonymous and non-synonymous single-nucleotide variants (SNVs) and insertions and deletions (indels) per area of coding genome sampled, after the removal of known and likely oncogenic driver events and germline SNPs. Tissue TMB is calculated based on variants with an allele frequency of ≥5%, and bTMB is calculated based on variants with an allele frequency of ≥0.5%.

Not Tested = not baited, not reported on test, or test preceded addition of biomarker or gene

Not Detected = baited but not detected on test

Detected = present (VAF% is not applicable)

VAF% = variant allele frequency percentage

 ${\sf Cannot\,Be\,Determined\,=\,Sample\,is\,not\,of\,sufficient\,data\,quality\,to\,confidently\,determine\,biomarker\,status}$ 

**BIOMARKER FINDINGS** 

#### BIOMARKER

# Blood Tumor Mutational Burden

RESULT 1 Muts/Mb

#### **POTENTIAL TREATMENT STRATEGIES**

On the basis of clinical evidence in NSCLC and HSNCC, increased bTMB may be associated with greater sensitivity to immunotherapeutic agents, including anti-PD-L1<sup>1-2</sup> and anti-PD-1<sup>3</sup> therapies. In NSCLC, multiple clinical trials have shown patients with higher bTMB derive clinical benefit from immune checkpoint inhibitors following single agent or combination treatments with either CTLA4 inhibitors or chemotherapy, with reported high bTMB cutpoints ranging from 6 to 16 Muts/Mb¹. In HNSCC, a Phase 3 trial showed that bTMB ≥16 Muts/Mb (approximate

equivalency ≥8 Muts/Mb as measured by this assay) was associated with improved survival from treatment with a PD-L1 inhibitor alone or in combination with a CTLA-4 inhibitor<sup>4</sup>.

#### **FREQUENCY & PROGNOSIS**

Average bTMB levels in solid tumors other than NSCLC have not been evaluated (cBioPortal, COSMIC, PubMed, Mar 2021)5-7. Published data investigating the prognostic implications of bTMB levels in breast cancer are limited (PubMed, Jul 2020). In a large study of patients with breast cancer, hypermutation was more frequently observed in metastatic tumors than in primary tumors8. In a study of 14,867 patients with breast cancer, high TMB was associated with older age and metastatic disease but was not significantly associated with PD-L1 positivity using the TMB cutoff of ≥10 Muts/Mb9. In estrogen receptorpositive breast cancer, increased TMB in tissue samples (>mean of 1.25 Muts/Mb) associated with shorter OS (HR=2.02) in an analysis of the TCGA data10

#### **FINDING SUMMARY**

Blood tumor mutational burden (bTMB, also known as mutation load) is a measure of the number of somatic protein-coding base substitution and insertion/deletion mutations from circulating tumor DNA in blood. TMB is affected by a variety of causes, including exposure to mutagens such as ultraviolet light in melanoma<sup>11-12</sup> and cigarette smoke in lung cancer<sup>13-14</sup>, treatment with temozolomide-based chemotherapy in glioma<sup>15-16</sup>, mutations in the proofreading domains of DNA polymerases encoded by the POLE and POLD1 genes<sup>17-21</sup>, and microsatellite instability (MSI)17,20-21. High bTMB levels were not detected in this sample. It is unclear whether the bTMB levels in this sample would be predicted to be associated with sensitivity to PD-1- or PD-L1-targeting immune checkpoint inhibitors, alone or in combination with other agents<sup>1-3</sup>. Depending on the clinical context, TMB testing of an alternate sample or by another methodology could be considered.

#### BIOMARKER

# **Tumor Fraction**

RESULT

Cannot Be Determined

#### POTENTIAL TREATMENT STRATEGIES

Specimens with high tumor fraction values have high circulating-tumor DNA (ctDNA) content, and thus high sensitivity for identifying genomic alterations. Such specimens are at low risk of false negative results. However, if tumor fraction is not detected as high, it does not exclude the presence of disease burden or compromise the confidence of reported alterations. Tumor fraction levels currently have limited implications for diagnosis, surveillance, or therapy and should not be overinterpreted or compared from one blood draw

to another. There are currently no targeted approaches to address specific tumor fraction levels. In the research setting, changes in tumor fraction estimates have been associated with treatment duration and clinical response and may be a useful indicator for future cancer management<sup>22-27</sup>.

#### **FREQUENCY & PROGNOSIS**

Detectible ctDNA levels have been reported in a variety of tumor types, with higher tumor fraction levels reported for patients with metastatic (Stage 4) tumors compared with patients with localized disease (Stages 1 to 3)<sup>28</sup>. Elevated tumor fraction levels have been reported to be associated with worse prognosis in a variety of cancer types, including pancreatic cancer<sup>29</sup>, Ewing sarcoma and osteosarcoma<sup>30</sup>, prostate cancer<sup>25</sup>, breast cancer<sup>31</sup>, leiomyosarcoma<sup>32</sup>, esophageal cancer<sup>33</sup>, colorectal cancer<sup>34</sup>, and gastrointestinal cancer<sup>35</sup>.

#### **FINDING SUMMARY**

Tumor fraction provides an estimate of the percentage of ctDNA present in a cell-free DNA (cfDNA) sample. The tumor fraction estimate for this sample is based on the observed level of aneuploid instability. The tumor fraction algorithm utilized for FoundationOne Liquid CDx uses the allele frequencies of approximately 1,000 singlenucleotide polymorphism (SNP) sites across the genome. Unlike the maximum somatic allele frequency (MSAF) method of estimating ctDNA content<sup>36</sup>, the tumor fraction metric does not take into account the allele frequency of individual variants but rather produces a more holistic estimate of ctDNA content using data from across the genome. The amount of ctDNA detected may correlate with disease burden and response to therapy  $^{37-38}$ . However, the tumor fraction estimate in this sample could not be determined with confidence.

**GENOMIC FINDINGS** 

**GENE** 

# FGF6

ALTERATION R62H

TRANSCRIPT ID NM\_020996

CODING SEQUENCE EFFECT

185G>A

#### **POTENTIAL TREATMENT STRATEGIES**

There are no targeted therapies that directly address genomic alterations in FGF6. Inhibitors of FGF receptors, however, are undergoing clinical trials in a number of different cancers. Limited data suggest that pan-FGFR inhibitors show activity in FGF amplified cancers; following treatment with a selective pan-FGFR inhibitor, a patient with head and neck squamous cell

carcinoma (HNSCC) and amplification of 11q13 (FGF3, FGF4, FGF19) and 12p13 (FGF6 and FGF23) experienced a radiologic CR<sup>39</sup>.

#### **FREQUENCY & PROGNOSIS**

Somatic alterations affecting FGF6 are infrequently documented, with the highest rates reported in penile cancer (4%), cutaneous melanoma (1-3%), stomach carcinoma (1-3%) and colorectal cancer (1%) (cBioPortal, COSMIC, 2021)5-7. Amplification of FGF6 has been frequently observed in testicular germ cell cancer (5%) and ovarian serous cystadenocarcinoma (5%), and in 2-6% of lower-grade gliomas, glioblastomas, sarcomas, breast invasive carcinomas, uterine carcinosarcomas, lung squamous cell carcinomas (SCC), head and neck SCC, pancreatic adenocarcinomas, and esophageal carcinomas (cBioPortal, 2021)5-6. FGF6 is colocalized with FGF23 and CCND2 at chromosomal locus 12p13 and has been reported

to be co-amplified with these genes in 1.3% of patients with breast cancer  $^{40}$ . FGF6 expression has been reported in 54% (14/26) of prostate cancer samples, which also frequently express FGFR4 $^{41}$ . FGF6 expression has also been observed in 71% (12/17) of patients with childhood acute lymphoblastic leukemia $^{42}$ .

#### **FINDING SUMMARY**

FGF6 (also known as HST-2) encodes a member of the fibroblast growth factor protein family and is hypothesized to play a role in muscle tissue regeneration<sup>43</sup> by signaling through FGFR<sub>4</sub>, and to a lesser extent FGFR<sub>1</sub> and FGFR<sub>2</sub><sup>44</sup>. FGF6 expression has been observed in several cancers<sup>41-42,45</sup> and was shown to be oncogenic in preclinical models<sup>45-46</sup>. FGF6 has been reported as amplified in cancer<sup>6</sup> and may be biologically relevant in this context<sup>47-48</sup>.



TUMOR TYPE
Breast cancer (NOS)

REPORT DATE 21 May 2021



ORDERED TEST # ORD-1095717-01

APPENDIX

Variants of Unknown Significance

**NOTE** One or more variants of unknown significance (VUS) were detected in this patient's tumor. These variants may not have been adequately characterized in the scientific literature at the time this report was issued, and/or the genomic context of these alterations makes their significance unclear. We choose to include them here in the event that they become clinically meaningful in the future.

 ATRX
 BARD1
 CBL
 DDR2

 R907Q
 E445K
 C404F
 E557D

 KEAP1
 KMT2A (MLL)
 MLL2
 PIK3C2B

 V369A
 A53V
 A765\_Q773del
 A858T

**RET** H54Y

**APPENDIX** 

Genes assayed in FoundationOne®Liquid CDx

FoundationOne Liquid CDx interrogates 324 genes, including 309 genes with complete exonic (coding) coverage and 15 genes with only select non-coding coverage (indicated with an \*); 75 genes (indicated in bold) are captured with increased sensitivity and have complete exonic (coding) coverage unless otherwise noted.

| ABL1<br>Exons 4-9          | ACVR1B                                       | AKT1<br>Exon 3              | AKT2                              | AKT3                                                     | ALK<br>Exons 20-29, Introns<br>18, 19           | ALOX12B                                                               | AMER1<br>(FAM123B)                          | APC             |
|----------------------------|----------------------------------------------|-----------------------------|-----------------------------------|----------------------------------------------------------|-------------------------------------------------|-----------------------------------------------------------------------|---------------------------------------------|-----------------|
| AR                         | <b>ARAF</b><br>Exons 4, 5, 7, 11, 13, 15, 16 | ARFRP1                      | ARID1A                            | ASXL1                                                    | ATM                                             | ATR                                                                   | ATRX                                        | AURKA           |
| AURKB                      | AXIN1                                        | AXL                         | BAP1                              | BARD1                                                    | BCL2                                            | BCL2L1                                                                | BCL2L2                                      | BCL6            |
| BCOR                       | BCORL1                                       | BCR*<br>Introns 8, 13, 14   | BRAF<br>Exons 11-18, Introns 7-10 | <b>BRCA1</b> Introns 2, 7, 8, 12, 16, 19, 20             | BRCA2<br>O Intron 2                             | BRD4                                                                  | BRIP1                                       | BTG1            |
| BTG2                       | BTK<br>Exons 2, 15                           | C11orf30<br>(EMSY)          | C17orf39<br>(GID4)                | CALR                                                     | CARD11                                          | CASP8                                                                 | CBFB                                        | CBL             |
| CCND1                      | CCND2                                        | CCND3                       | CCNE1                             | CD22                                                     | CD70                                            | CD74*<br>Introns 6-8                                                  | CD79A                                       | CD79B           |
| CD274<br>(PD-L1)           | CDC73                                        | CDH1                        | CDK12                             | CDK4                                                     | CDK6                                            | CDK8                                                                  | CDKN1A                                      | CDKN1B          |
| CDKN2A                     | CDKN2B                                       | CDKN2C                      | СЕВРА                             | CHEK1                                                    | CHEK2                                           | CIC                                                                   | CREBBP                                      | CRKL            |
| CSF1R                      | CSF3R                                        | CTCF                        | CTNNA1                            | CTNNB1<br>Exon 3                                         | CUL3                                            | CUL4A                                                                 | CXCR4                                       | CYP17A1         |
| DAXX                       | DDR1                                         | <b>DDR2</b> Exons 5, 17, 18 | DIS3                              | DNMT3A                                                   | DOT1L                                           | EED                                                                   | EGFR<br>Introns 7, 15, 24-27                | EP300           |
| ЕРНАЗ                      | ЕРНВ1                                        | ЕРНВ4                       | ERBB2                             | ERBB3<br>Exons 3, 6, 7, 8, 10, 12,<br>20, 21, 23, 24, 25 | ERBB4                                           | ERCC4                                                                 | ERG                                         | ERRFI1          |
| ESR1<br>Exons 4-8          | ETV4*<br>Intron 8                            | ETV5*<br>Introns 6, 7       | ETV6*<br>Introns 5, 6             | EWSR1*<br>Introns 7-13                                   | <b>EZH2</b><br>Exons 4, 16, 17, 18              | EZR*<br>Introns 9-11                                                  | FAM46C                                      | FANCA           |
| FANCC                      | FANCG                                        | FANCL                       | FAS                               | FBXW7                                                    | FGF10                                           | FGF12                                                                 | FGF14                                       | FGF19           |
| FGF23                      | FGF3                                         | FGF4                        | FGF6                              | FGFR1<br>Introns 1, 5, Intron 17                         | FGFR2<br>Intron 1, Intron 17                    | FGFR3 Exons 7, 9 (alternative designation exon 10), 14, 18, Intron 17 | FGFR4                                       | FH              |
| FLCN                       | FLT1                                         | FLT3<br>Exons 14, 15, 20    | FOXL2                             | FUBP1                                                    | GABRA6                                          | GATA3                                                                 | GATA4                                       | GATA6           |
| <b>GNA11</b><br>Exons 4, 5 | GNA13                                        | GNAQ<br>Exons 4, 5          | GNAS<br>Exons 1, 8                | GRM3                                                     | GSK3B                                           | НЗГЗА                                                                 | HDAC1                                       | HGF             |
| HNF1A                      | HRAS<br>Exons 2, 3                           | HSD3B1                      | ID3                               | IDH1<br>Exon 4                                           | IDH2<br>Exon 4                                  | IGF1R                                                                 | IKBKE                                       | IKZF1           |
| INPP4B                     | IRF2                                         | IRF4                        | IRS2                              | JAK1                                                     | JAK2<br>Exon 14                                 | <b>JAK3</b><br>Exons 5, 11, 12, 13, 15, 16                            | JUN                                         | KDM5A           |
| KDM5C                      | KDM6A                                        | KDR                         | KEAP1                             | KEL                                                      | <b>KIT Exons 8, 9, 11, 12, 13, 17</b> Intron 16 | KLHL6<br>7,                                                           | KMT2A<br>(MLL) Introns 6, 8-11,<br>Intron 7 | KMT2D<br>(MLL2) |

APPENDIX

Genes assayed in FoundationOne®Liquid CDx

FoundationOne Liquid CDx interrogates 324 genes, including 309 genes with complete exonic (coding) coverage and 15 genes with only select non-coding coverage (indicated with an \*); 75 genes (indicated in bold) are captured with increased sensitivity and have complete exonic (coding) coverage unless otherwise noted.

| KRAS                      | LTK                                                    | LYN                                                  | MAF                                                          | MAP2K1<br>(MEK1) Exons 2, 3                                   | MAP2K2<br>(MEK2) Exons 2-4, 6, | MAP2K4<br>7         | МАРЗК1                | МАРЗК1З                                                      |
|---------------------------|--------------------------------------------------------|------------------------------------------------------|--------------------------------------------------------------|---------------------------------------------------------------|--------------------------------|---------------------|-----------------------|--------------------------------------------------------------|
| МАРК1                     | MCL1                                                   | MDM2                                                 | MDM4                                                         | MED12                                                         | MEF2B                          | MEN1                | MERTK                 | MET                                                          |
| MITF                      | MKNK1                                                  | MLH1                                                 | MPL<br>Exon 10                                               | MRE11A                                                        | MSH2<br>Intron 5               | MSH3                | MSH6                  | MST1R                                                        |
| МТАР                      | MTOR<br>Exons 19, 30, 39, 40,<br>43-45, 47, 48, 53, 56 | МИТҮН                                                | MYB*<br>Intron 14                                            | MYC<br>Intron 1                                               | MYCL<br>(MYCL1)                | MYCN                | MYD88<br>Exon 4       | NBN                                                          |
| NF1                       | NF2                                                    | NFE2L2                                               | NFKBIA                                                       | NKX2-1                                                        | NOTCH1                         | NOTCH2<br>Intron 26 | <b>NOTCH3</b>         | <b>NPM1</b><br>Exons 4-6, 8, 10                              |
| NRAS<br>Exons 2, 3        | NSD3<br>(WHSC1L1)                                      | NT5C2                                                | <b>NTRK1</b><br>Exons 14, 15, Introns<br>8-11                | NTRK2<br>Intron 12                                            | NTRK3<br>Exons 16, 17          | NUTM1*<br>Intron 1  | P2RY8                 | PALB2                                                        |
| PARK2                     | PARP1                                                  | PARP2                                                | PARP3                                                        | PAX5                                                          | PBRM1                          | PDCD1<br>(PD-1)     | PDCD1LG2<br>(PD-L2)   | <b>PDGFRA</b><br>Exons 12, 18, Introns 7,<br>9, 11           |
| PDGFRB<br>Exons 12-21, 23 | PDK1                                                   | PIK3C2B                                              | PIK3C2G                                                      | PIK3CA<br>Exons 2, 3, 5-8, 10, 14,<br>19, 21 (Coding Exons 1, | PIK3CB                         | PIK3R1              | PIM1                  | PMS2                                                         |
| POLD1                     | POLE                                                   | PPARG                                                | PPP2R1A                                                      | 2, 4-7, 9, 13, 18, 20)<br>PPP2R2A                             | PRDM1                          | PRKAR1A             | PRKCI                 | РТСН1                                                        |
| PTEN                      | PTPN11                                                 | PTPRO                                                | QKI                                                          | RAC1                                                          | RAD21                          | RAD51               | RAD51B                | RAD51C                                                       |
| RAD51D                    | RAD52                                                  | RAD54L                                               | <b>RAF1</b><br>Exons 3, 4, 6, 7, 10, 14, 15, 17, Introns 4-8 | RARA<br>Intron 2                                              | RB1                            | RBM10               | REL                   | <b>RET</b><br>Introns 7, 8, Exons 11,<br>13-16, Introns 9-11 |
| RICTOR                    | RNF43                                                  | <b>ROS1</b><br>Exons 31, 36-38, 40,<br>Introns 31-35 | RPTOR                                                        | RSPO2*<br>Intron 1                                            | SDC4*<br>Intron 2              | SDHA                | SDHB                  | SDHC                                                         |
| SDHD                      | SETD2                                                  | SF3B1                                                | SGK1                                                         | SLC34A2*<br>Intron 4                                          | SMAD2                          | SMAD4               | SMARCA4               | SMARCB1                                                      |
| SMO                       | SNCAIP                                                 | SOCS1                                                | SOX2                                                         | SOX9                                                          | SPEN                           | SPOP                | SRC                   | STAG2                                                        |
| STAT3                     | STK11                                                  | SUFU                                                 | SYK                                                          | ТВХЗ                                                          | TEK                            | TERC*<br>ncRNA      | <b>TERT*</b> Promoter | TET2                                                         |
| TGFBR2                    | TIPARP                                                 | TMPRSS2*<br>Introns 1-3                              | TNFAIP3                                                      | TNFRSF14                                                      | TP53                           | TSC1                | TSC2                  | TYRO3                                                        |
| U2AF1                     | VEGFA                                                  | VHL                                                  | WHSC1                                                        | WTI                                                           | XPO1                           | XRCC2               | ZNF217                | ZNF703                                                       |

ADDITIONAL ASSAYS: FOR THE DETECTION OF SELECT CANCER BIOMARKERS

Microsatellite (MS) status

Blood Tumor Mutational Burden (bTMB)

**Tumor Fraction** 

**APPENDIX** 

About FoundationOne®Liquid CDx

FoundationOne Liquid CDx fulfills the requirements of the European Directive 98/79 EC for in vitro diagnostic medical devices and is registered as a CE-IVD product by Foundation Medicine's EU Authorized Representative, Qarad b.v.b.a, Cipalstraat 3, 2440 Geel, Belgium. The CE-IVD regulatory status of FoundationOne Liquid CDx is applicable in countries that accept and/or recognize the CE mark.





#### **ABOUT FOUNDATIONONE LIQUID CDX**

FoundationOne Liquid CDx was developed and its performance characteristics determined by Foundation Medicine, Inc. (Foundation Medicine). FoundationOne Liquid CDx may be used for clinical purposes and should not be regarded as purely investigational or for research only. Foundation Medicine's clinical reference laboratories are qualified to perform highcomplexity clinical testing.

Please refer to technical information for performance specification details.

#### **INTENDED USE**

FoundationOne Liquid CDx is a next generation sequencing based in vitro diagnostic device that analyzes 324 genes. Substitutions and insertion and deletion alterations (indels) are reported in 311 genes, copy number alterations (CNAs) are reported in 310 genes, and gene rearrangements are reported in 324 genes. The test also detects the genomic signatures blood tumor mutational burden (bTMB), microsatellite instability (MSI), and tumor fraction. FoundationOne Liquid CDx utilizes circulating cell-free DNA (cfDNA) isolated from plasma derived from the anti-coagulated peripheral whole blood of cancer patients. The test is intended to be used as a companion diagnostic to identify patients who may benefit from treatment with targeted therapies in accordance with the approved therapeutic product labeling. Additionally, FoundationOne Liquid CDx is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for patients with malignant neoplasms.

### **TEST PRINCIPLES**

The FoundationOne Liquid CDx assay is performed exclusively as a laboratory service using circulating cell-free DNA (cfDNA) isolated from plasma derived from anti-coagulated peripheral whole blood from patients with solid malignant neoplasms. The assay employs a single DNA extraction method to obtain cfDNA from plasma from whole blood. Extracted

cfDNA undergoes whole-genome shotgun library construction and hybridization-based capture of 324 cancer-related genes including coding exons and select introns of 309 genes, as well as only select intronic regions or non-coding regions of 15 genes. Hybrid-capture selected libraries are sequenced with deep coverage using the NovaSeq® 6000 platform. Sequence data are processed using a customized analysis pipeline designed to accurately detect genomic alterations, including base substitutions, indels, select copy number variants, and select genomic rearrangements. Substitutions and insertion and deletion alterations (indels) are reported in 311 genes, copy number alterations (CNAs) are reported in 310 genes, and gene rearrangements are reported in 324 genes. The assay also detects select genomic rearrangements, select copy number alterations, tumor fraction, and genomic signatures including MSI and bTMB. A subset of targeted regions in 75 genes is baited for increased sensitivity.

#### THE REPORT

Incorporates analyses of peer-reviewed studies and other publicly available information identified by Foundation Medicine; these analyses and information may include associations between a molecular alteration (or lack of alteration) and one or more drugs with potential clinical benefit (or potential lack of clinical benefit), including drug candidates that are being studied in clinical research. Note: A finding of biomarker alteration does not necessarily indicate pharmacologic effectiveness (or lack thereof) of any drug or treatment regimen; a finding of no biomarker alteration does not necessarily indicate lack of pharmacologic effectiveness (or effectiveness) of any drug or treatment regimen.

#### **QUALIFIED ALTERATION CALLS** (EQUIVOCAL)

All equivocal calls, regardless of alteration type, imply that there is adequate evidence to call the alteration with confidence. However, the repeatability of equivocal calls may be lower than non-equivocal calls.

#### **RANKING OF ALTERATIONS AND THERAPIES**

Biomarker and Genomic Findings Therapies are ranked based on the following criteria: Therapies with clinical benefit in patient's tumor type (ranked alphabetically within each NCCN category) followed by therapies with clinical benefit in other tumor type (ranked alphabetically within each NCCN category).

Clinical Trials

Pediatric trial qualification → Geographical proximity → Later trial phase.

#### **LIMITATIONS**

- 1. For in vitro diagnostic use.
- 2. For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- 3. A negative result does not rule out the presence of a mutation below the limits of detection of the assay. Patients for whom no companion diagnostic alterations are detected should be considered for confirmation with an appropriately validated tumor tissue test, if available.
- 4. The FoundationOne Liquid CDx assay does not detect heterozygous deletions.
- **5**. The test is not intended to provide information on cancer predisposition.
- 6. Performance has not been validated for cfDNA input below the specified minimum input.
- 7. Tissue TMB and blood TMB (bTMB) are estimated from the number of synonymous and nonsynonymous single-nucleotide variants (SNVs) and insertions and deletions (indels) per area of coding genome sampled, after the removal of known and likely oncogenic driver events and germline SNPs. Tissue TMB is calculated based on variants with an allele frequency of ≥5%, and bTMB is calculated based on variants with an allele frequency of ≥0.5%.
- 8. Tumor fraction is the percentage of circulatingtumor DNA (ctDNA) present in a cell-free DNA (cfDNA) sample. The tumor fraction estimate is computationally derived from observed aneuploid instability in the sample.
- 9. Microsatellite instability (MSI) is a condition of genetic hypermutability that generates excessive amounts of short insertion/deletion mutations in the tumor genome; it generally occurs at microsatellite DNA sequences and is caused by a deficiency in DNA mismatch repair (MMR) in the tumor. The MSI algorithm is based on genome wide analysis of 1765 microsatellite loci and not based on the 5 or 7 MSI loci described in current clinical practice guidelines for solid tissue testing.
- 10. Genomic findings from circulating cell-free DNA (cfDNA) may originate from circulating tumor DNA fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP). Genes with alterations that may be derived from CHIP include, but are not limited

APPENDIX

About FoundationOne®Liquid CDx

to: ASXL1, ATM, CBL, CHEK2, DNMT3A, JAK2, KMT2D (MLL2), MPL, MYD88, SF3B1, TET2, TP53, and U2AF1.

- 11. Alterations reported may include somatic (not inherited) or germline (inherited) alterations; however, the test does not distinguish between germline and somatic alterations. If a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- 12. The test is not intended to replace germline testing or to provide information about cancer predisposition.

#### VARIANTS TO CONSIDER FOR FOLLOW-UP GERMLINE TESTING

The variants indicated for consideration of followup germline testing are 1) limited to reportable short variants with a protein effect listed in the ClinVar genomic database (Landrum et al., 2018; 29165669) as Pathogenic, Pathogenic/Likely Pathogenic, or Likely Pathogenic (by an expert panel or multiple submitters), 2) associated with hereditary cancer-predisposing disorder(s), 3) detected at an allele frequency of >30%, and 4) in select genes reported by the ESMO Precision Medicine Working Group (Mandelker et al., 2019; 31050713) to have a greater than 10% probability of germline origin if identified during tumor sequencing. The selected genes are ATM, BAP1, BRCA1, BRCA2, BRIP1, CHEK2, FH, FLCN, MLH1, MSH2, MSH6, MUTYH, PALB2, PMS2, POLE, RAD51C, RAD51D, RET, SDHA, SDHB, SDHC, SDHD, TSC2, and VHL, and are not inclusive of all cancer susceptibility genes. The content in this report should not substitute for genetic counseling or follow-up germline testing, which is needed to distinguish whether a finding in this patient's tumor sequencing is germline or somatic. Interpretation should be based on clinical context.

# NATIONAL COMPREHENSIVE CANCER NETWORK\* (NCCN\*) CATEGORIZATION

Biomarker and genomic findings detected may be associated with certain entries within the NCCN Drugs & Biologics Compendium® (NCCN Compendium®) (www.nccn.org). The NCCN Categories of Evidence and Consensus indicated reflect the highest possible category for a given therapy in association with each biomarker or genomic finding. Please note, however, that the accuracy and applicability of these NCCN categories within a report may be impacted by the patient's clinical history, additional biomarker information, age, and/or co-occurring alterations. For additional information on the NCCN categories, please refer to the NCCN

Compendium®. Referenced with permission from the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®). © National Comprehensive Cancer Network, Inc. 2021. All rights reserved. To view the most recent and complete version of the guidelines, go online to NCCN.org. NCCN makes no warranties of any kind whatsoever regarding their content, use or application and disclaims any responsibility for their application or use in any way.

#### **LEVEL OF EVIDENCE NOT PROVIDED**

Drugs with potential clinical benefit (or potential lack of clinical benefit) are not evaluated for source or level of published evidence.

#### **NO GUARANTEE OF CLINICAL BENEFIT**

This report makes no promises or guarantees that a particular drug will be effective in the treatment of disease in any patient. This report also makes no promises or guarantees that a drug with potential lack of clinical benefit will in fact provide no clinical benefit.

#### **NO GUARANTEE OF REIMBURSEMENT**

Foundation Medicine makes no promises or guarantees that a healthcare provider, insurer or other third party payor, whether private or governmental, will reimburse a patient for the cost of FoundationOne Liquid CDx.

# TREATMENT DECISIONS ARE THE RESPONSIBILITY OF PHYSICIAN

Drugs referenced in this Report may not be suitable for a particular patient. The selection of any, all or none of the drugs associated with potential clinical benefit (or potential lack of clinical benefit) resides entirely within the discretion of the treating physician. Indeed, the information in this Report must be considered in conjunction with all other relevant information regarding a particular patient, before the patient's treating physician recommends a course of treatment. Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and and patient preferences, in accordance with the standard of care in a given community. A treating physician's decisions should not be based on a single test, such as this test or the information contained in this report.

Certain sample of variant characteristics may result in reduced sensitivity. These include: low sample quality, deletions and insertions >4obp, or repetitive/high homology sequences. FoundationOne Liquid CDx is performed using cell-free DNA, and as such germline events may not be reported.

#### **SELECT ABBREVIATIONS**

| ABBREVIATION | DEFINITION                  |  |  |  |
|--------------|-----------------------------|--|--|--|
| CR           | Complete response           |  |  |  |
| DCR          | Disease control rate        |  |  |  |
| DNMT         | DNA methyltransferase       |  |  |  |
| HR           | Hazard ratio                |  |  |  |
| ITD          | Internal tandem duplication |  |  |  |
| MMR          | Mismatch repair             |  |  |  |
| Muts/Mb      | Mutations per megabase      |  |  |  |
| NOS          | Not otherwise specified     |  |  |  |
| ORR          | Objective response rate     |  |  |  |
| os           | Overall survival            |  |  |  |
| PD           | Progressive disease         |  |  |  |
| PFS          | Progression-free survival   |  |  |  |
| PR           | Partial response            |  |  |  |
| SD           | Stable disease              |  |  |  |
| TKI          | Tyrosine kinase inhibitor   |  |  |  |

MR Suite Version 4.0.0

(2013) pmid: 23636398

20. Nature (2012) pmid: 22810696

25568919

pmid: 30923679

#### ORDERED TEST # ORD-1095717-01



- 1. Gandara DR, et al. Nat. Med. (2018) pmid: 30082870
- 2. Wang Z, et al. JAMA Oncol (2019) pmid: 30816954
- Aggarwal C, et al. Clin. Cancer Res. (2020) pmid: 32102950
- 4. Li et al., 2020; ASCO Abstract 6511
- 5. Cerami E, et al. Cancer Discov (2012) pmid: 22588877
- 6. Gao J, et al. Sci Signal (2013) pmid: 23550210
- 7. Tate JG, et al. Nucleic Acids Res. (2019) pmid: 30371878
- 8. Barroso-Sousa R, et al. Ann. Oncol. (2020) pmid: 32067680
- 9. Chumsri S, et al. J Natl Compr Canc Netw (2020) pmid: 32380464
- 10. Haricharan S, et al. Breast Cancer Res. Treat. (2014) pmid: 24839032
- 11. Pfeifer GP, et al. Mutat. Res. (2005) pmid: 15748635
- 12. Hill VK, et al. Annu Rev Genomics Hum Genet (2013) pmid: 23875803
- 13. Pfeifer GP, et al. Oncogene (2002) pmid: 12379884
- 14. Rizvi NA, et al. Science (2015) pmid: 25765070
- 15. Johnson BE, et al. Science (2014) pmid: 24336570 16. Choi S, et al. Neuro-oncology (2018) pmid: 29452419
- 23. Raja R, et al. Clin. Cancer Res. (2018) pmid: 30093454

17. Cancer Genome Atlas Research Network, et al. Nature

19. Heitzer E, et al. Curr. Opin. Genet. Dev. (2014) pmid:

18. Briggs S, et al. J. Pathol. (2013) pmid: 23447401

21. Roberts SA, et al. Nat. Rev. Cancer (2014) pmid:

22. Bronkhorst AJ, et al. Biomol Detect Quantif (2019)

- 24. Hrebien S, et al. Ann. Oncol. (2019) pmid: 30860573
- 25. Choudhury AD, et al. JCI Insight (2018) pmid: 30385733
- 26. Goodall J, et al. Cancer Discov (2017) pmid: 28450425
- 27. Goldberg SB, et al. Clin. Cancer Res. (2018) pmid:
- 28. Bettegowda C, et al. Sci Transl Med (2014) pmid: 24553385
- 29. Lapin M, et al. J Transl Med (2018) pmid: 30400802
- 30. Shulman DS, et al. Br. J. Cancer (2018) pmid: 30131550
- 31. Stover DG, et al. J. Clin. Oncol. (2018) pmid: 29298117

- 32. Hemming ML, et al. JCO Precis Oncol (2019) pmid: 30793095
- 33. Egyud M, et al. Ann. Thorac. Surg. (2019) pmid: 31059681
- 34. Fan G, et al. PLoS ONE (2017) pmid: 28187169
- 35. Vu et al., 2020; DOI: 10.1200/PO.19.00204
- 36. Li G, et al. J Gastrointest Oncol (2019) pmid: 31602320
- 37. Zhang EW, et al. Cancer (2020) pmid: 32757294
- 38. Butler TM, et al. Cold Spring Harb Mol Case Stud (2019) pmid: 30833418
- 39. Dumbrava et al., 2018; doi/full/10.1200/PO.18.00100
- 40. Parish A, et al. Cell Cycle (2015) pmid: 25950492
- 41. Ropiquet F, et al. Cancer Res. (2000) pmid: 10945637
- 42. Niini T, et al. Leukemia (2002) pmid: 12399964
- 43. Neuhaus P, et al. Mol. Cell. Biol. (2003) pmid: 12917328
- 44. Ornitz DM, et al. J. Biol. Chem. (1996) pmid: 8663044
- 45. Iida S, et al. Oncogene (1992) pmid: 1549352
- **46.** Marics I, et al. Oncogene (1989) pmid: 2649847
- 47. Zack TI, et al. Nat. Genet. (2013) pmid: 24071852
- 48. Beroukhim R, et al. Nature (2010) pmid: 20164920