

# ANA LISI

Patologia molecolare



VIOLLIER

## Cancer Panels

### Oncomine™ Comprehensive Assay v3 DNA, Thermo Fisher – 145 geni

Mutazioni hotspot

AKT1, AKT2, AKT3, ALK, AR, ARAF, AXL, BRAF, BTK, CBL, CCND1, CDK4, CDK6, CHEK2, CSF1R, CTNNB1, DDR2, EGFR, ERBB2 (=HER2), ERBB3, ERBB4, ERCC2, ESR1, EZH2, FGFR1, FGFR2, FGFR3, FGFR4, FLT3, FOXL2, GATA2, GNA11, GNAQ, GNAS, H3F3A, HNF1A, HRAS, IDH1, IDH2, JAK1, JAK2, JAK3, KDR, KIT, KNSTRN, KRAS, MAGOH, MAP2K1 (=MEK1), MAP2K2 (=MEK2), MAP2K4, MAPK1, MAX, MDM2, MDM4, MED12, MET, MTOR, MYC, MYCL, MYCN, MYD88, NFE2L2, NRAS, NTRK1, NTRK2, NTRK3, PDGFR, PDGFRB, PIK3CA, PIK3CB, PPP2R1A, PTPN11, RAC1, RAF1, RET, RHEB, RHOA, ROS1, SF3B1, SMAD4, SMO, SPOP, SRC, STAT3, TERT, TOP1, U2AF1, XPO1

Sequenziamento completo del gene

ARID1A, ATM, ATR, ATRX, BAP1, BRCA1, BRCA2, CDK12, CDKN1B, CDKN2A, CDKN2B, CHEK1, CREBBP, FANCA, FANCD2, FANCI, FBXW7, MLH1, MRE11, MSH2, MSH6, NBN, NF1, NF2, NOTCH1, NOTCH2, NOTCH3, PALB2, PIK3R1, PMS2, POLE, PTCH1, PTEN, RAD50, RAD51, RAD51B, RAD51C, RAD51D, RB1, RNF43, SETD2, SLX4, SMARCA4, SMARCB1, STK11, TP53, TSC1, TSC2

Alterazioni del numero di copie (CNVs)

AKT1, AKT2, AKT3, ALK, AR, AXL, BRAF, CCND1, CCND2, CCND3, CCNE1, CDK2, CDK4, CDK6, EGFR, ERBB2, ESR1, FGF19, FGF3, FGFR1, FGFR2, FGFR3, FGFR4, FLT3, IGF1R, KIT, KRAS, MDM2, MDM4, MET, MYC, MYCL, MYCN, NTRK1, NTRK2, NTRK3, PDGFR, PDGFRB, PIK3CA, PIK3CB, PPARG, RICTOR, TERT

---

### Oncomine™ Comprehensive Assay RNA, Thermo Fisher – 51 partner di fusione

Fusioni di geni

AKT2, ALK, AR, AXL, BRAF, BRCA1, BRCA2, CDKN2A, EGFR, ERBB2, ERBB4, ERG, ESR1, ETV1, ETV4, ETV5, FGFR1, FGFR2, FGFR3, FGR, FLT3, JAK2, KRAS, MDM4, MET, MYB, MYBL1, NF1, NOTCH1, NOTCH4, NRG1, NTRK1, NTRK2, NTRK3, NUTM1, PDGFR, PDGFRB, PIK3CA, PPARG, PRKACA, PRKACB, PTEN, RAD51B, RAF1, RB1, RELA, RET, ROS1, RSPO2, RSPO3, TERT

---

### Oncomine™ Precision Assay DNA, Thermo Fisher – 46 geni

Mutazioni hotspot

AKT1, AKT2, AKT3, ALK, AR, ARAF, BRAF, CDK4, CDKN2A, CHEK2, CTNNB1, EGFR, ERBB2 (=HER2), ERBB3, ERBB4, ESR1, FGFR1, FGFR2, FGFR3, FGFR4, FLT3, GNA11, GNAQ, GNAS, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1 (=MEK1), MAP2K2 (= MEK2), MET, MTOR, NRAS, NTRK1, NTRK2, NTRK3, PDGFR, PIK3CA, PTEN, RAF1, RET, ROS1, SMO, TP53

Alterazioni del numero di copie (CNVs)

ALK, AR, CD274, CDKN2A, EGFR, ERBB2, ERBB3, FGFR1, FGFR2, FGFR3, KRAS, MET, PIK3CA, PTEN

## **Oncomine™ Focus Assay RNA, Thermo Fisher – 23 partner di fusione**

Fusioni di geni

ABL1, ALK, AKT3, AXL, BRAF, EGFR, ERBB2, ERG, ETV1, ETV4, ETV5, FGFR1, FGFR2, FGFR3, MET, NTRK1, NTRK2, NTRK3, PDGFRA, PPARG, RAF1, RET, ROS1

---

### **NGS Carcinoma colorettale**

Mutazioni hotspot

KRAS, NRAS, BRAF, PIK3CA

---

### **NGS Carcinoma polmonare**

Mutazioni hotspot

EGFR, KRAS, BRAF, HER2, MET

Fusioni di geni

ALK1, ROS1, RET, NTRK1, NTRK2, NTRK3, MET Exon-14-Skipping

---

### **NGS melanoma**

Mutazioni hotspot

BRAF, KIT, NRAS, HRAS

---

### **NGS tumore stromale gastrointestinale (GIST)**

Mutazioni hotspot

KIT, PDGFRA

---

### **NGS Carcinoma della tiroide**

Mutazioni hotspot

AKT1, BRAF, CTNNB1, HRAS, KRAS, NRAS, PIK3CA, PTEN, RET, TERT, TP53

Fusioni di geni

RET, PPARG, NTRK1, NTRK2, NTRK3

---

### **NGS Carcinoma dell'endometrio**

Sequenziamento completo del gene

POLE, TP53, MLH1, MSH2, MSH6, PMS2

---

### **NGS Carcinoma uroteliale**

Mutazioni hotspot e fusioni

FGFR1, FGFR2, FGFR3, FGFR4

Sequenziamento completo del gene

TP53

### **NGS Carcinoma ovarico**

TruSight Oncology 500 HRD powered by Myriad, Illumina

Mutazioni in geni associati alla BRCAness incl. BRCA1, BRCA2, PALB2

Valutazione del Genomic Instability Score (GIS)

---

### **NGS Carcinoma mammario pannello grande**

TruSight Oncology 500, Illumina

Mutazioni in geni associati alla BRCAness incl. BRCA1, BRCA2, PALB2

Incl. PIK3CA, AKT1, PTEN, ESR1

---

### **NGS Carcinoma mammario pannello piccolo**

Oncomine™ Comprehensive Assay v3 DNA, Thermo Fisher

PIK3CA, AKT1, PTEN, ESR1

---

### **Biopsia liquida Carcinoma mammario**

Oncomine™ Precision Assay GX, Thermo Fisher

ESR1, PIK3CA, AKT1

---

### **NGS Carcinoma prostatico**

TruSight Oncology 500, Illumina

Mutazioni in geni associati alla BRCAness incl. BRCA1, BRCA2, ATM

Incl. analisi dell'instabilità microsatellitare e CDK12

---

Analisi supplementari:

**BRCA1, BRCA2** Sequenziamento completo del gene

**Genomic Instability Score (GIS)** TruSight Oncology 500 HRD powered by Myriad

**Instabilità dei microsatelliti (MSI)** Analisi della lunghezza di segmento

**Metilazione del promotore di MLH1** Analisi della lunghezza di segmento

**Carico mutazionale del tumore (TMB)** TruSight Oncology 500, Illumina

Presso Viollier tutte le analisi elencate sono accreditate secondo la norma

SN EN ISO 15189.

### **Informazioni**

Dr. rer. nat. Henriette Kurth, Specialista FAMH in medicina di laboratorio, Responsabile Biologia molecolare

Dr. med. Katharina Marston, FMH patologia, patologia molecolare, Resp. medicina patologia molecolare

Dr. med. Sophie Diebold Berger, FMH patologia, citopatologia, Responsabile Viollier Genève SA Patologia

PD Dr. med. Andreas Zettl, FMH patologia, candidato patologia molecolare, Responsabile Patologia

viollier.ch