## **Biofocus' Molecular Diagnostic Panel**





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- Molecular detection of infectious diseases
- Human & veterinary hereditary diseases / genetic predisposition
- Molecular Oncology

# **Aims of Molecular Oncology**



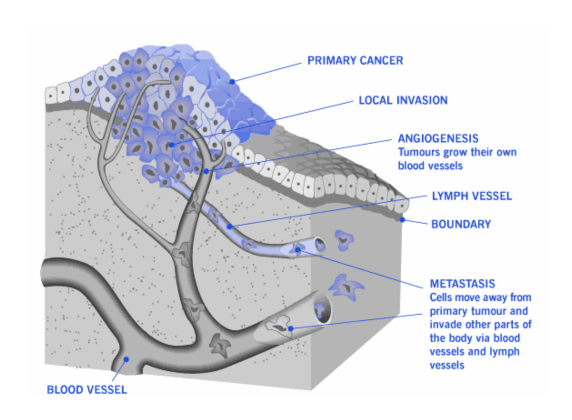
Detection von Circulating Tumor Cells in blood
 (CTCs) → "residual disease"

 Determination of "drug target"-genes and resistance-markes

→ Goal: personalized therapy







#### CTCs:

**rare:** 100 – 1000 per ml blood

sparely: 1 CTC in 10^6 to 10^7 WBCs

**Isolation of CTCs is challenging** 



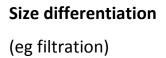
### Isolation of CTCs from blood by positive selection

#### blood sample

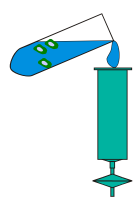
#### **CTC** isolation

Molecular characterisation



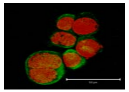










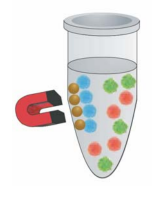


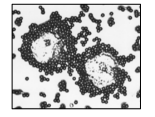




Immuno absorption

(eg. magnetic beads)







#### Molecular identification of captured cells as CTCs

- Isolation of RNA from captured CTCs
- Differential Gene expression CTCs ⇔ Blood
- Molecular Tumor markers by quantitative real-time PCR:
  - Cytokeratins (CK19, CK20)
  - Cell cycle genes (c-myc, erbb2, telomerase, survivin)
  - tissue specific genes: PSA (prostate), G250 (renal), MART (melanoma)



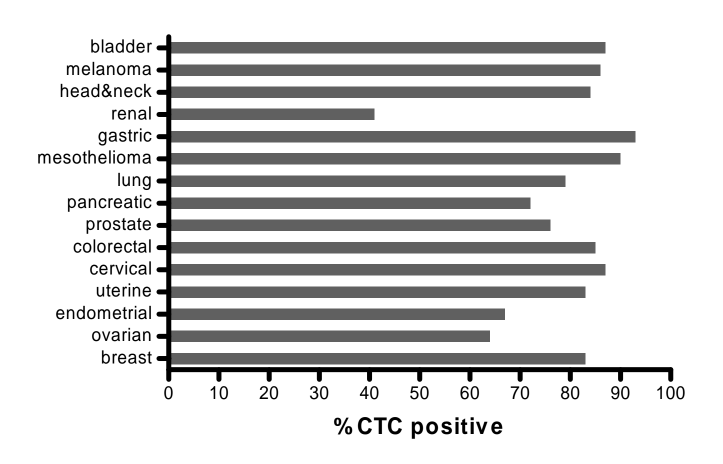


■ Four-Marker assay; Carcinoma (e.g. CK19, ERBB2, C-MYC, Telomerase):

	≥1 Marker positive	≥ 2 Marker positive
	positive	positive
Normal-	3/70	1/70
patients n = 70	4.3 %	1.4 %
Tumor CA Patients n = 200	159/200 80 %	121/200 60 %

#### **Detection rate of CTCs**





→ Average ca. 80 % in advanced tumors

#### **Molecular characterization of CTCs**



→ Gene expression analysis in CTCs for prediction of therapy resistance:

#### **Drug Metabolizing Genes**

Activation, Degradation, Detoxification

#### "Drug-Target" Genes

Cellular function inhibited by the drug



#### **Genetic factors of Chemo-Resistance**

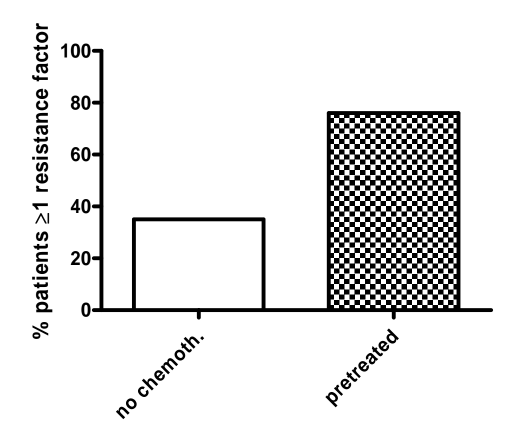
#### ■ Different drug-targets and metabolizing genes depending on drug

Drugs	Target	Resistance
Anthracyclines	Topoisomerase II	Topo II downregulation
Irinotecan	Topoisomerase I	Topo I downregulation
Platinum comp.	DNA	induction of ERCC1 repair
Nitroso-Ureas	DNA	induction of MGMT repair
Methotrexate	Folate metabolism	overexpression of DHFR
5-Fluorouracil	Nucleotide/DNA synthesis	overexpression of TS / DPD



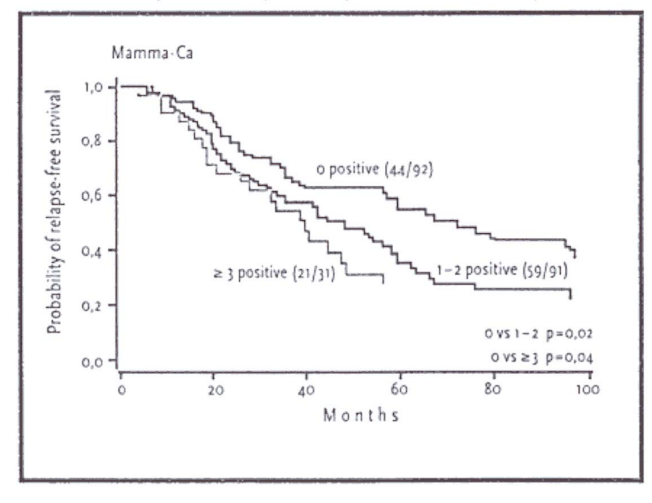
#### Multi-resistance in chemo-treated vs. non-treated patients

multidrug resistance factors: MDR1, MRP1, GST/GCS



#### Impact of resistance factors on relapse-free-survival

GST-π; γ-GCS; MDR1 (DNA; RNA); MRP; DHFR; TS; bcl-2/bax



# **Therapy: Prediction and Outcome**



Tumor type	Therapy	Test-Result	clinical outcome
Mamma	Mitoxantron	resistent	Progress
Colon	5-FU	resistent	Progress
Stomach	Mitomycin + 5-FU + Cyclophpos.	Mitomycin: resistent Cyclophpos.: resistent 5-FU: intermediate	Progress / death
Ovar	Gemcitabine + Cyclophpos.	Gemcitabine: resistent Cyclophpos: sensitiv	Progress / death
Mamma	MTX + Cisplatin	MTX: resistent Cisplatin.: resistent	Partial response
Colon	Oxaliplatin	sensitiv	Partial response
Mamma	MTX + Gemcitabin	MTX: sensitiv Gemcitabin: sensitiv	Partial response
Mamma	MTX + 5-FU	MTX: sensitiv 5-FU: intermediate	Partial response
Mamma	Herceptin	sensitiv	Partial response
Pancreas	Herceptin	sensitiv	Partial response
Thymoma	Epirubicin + Mitomycin + 5-FU	Mitomycin: sensitiv 5-FU: sensitiv Epirubicin: intermediate	Complete response
Melanoma	Sorafenib	sensitiv	Complete response

#### Limitations of the prediction model



#### **Resistance** ⇔ **Sensitivity**:

- Generally it is easier to predict resistance than response
- Focus on major resistance pathways only

#### **Alternative Therapies:**

Clinical response is observed despite positive resistance marker

→ Modulation of the resistance genes by alternative agents





#### Alternative agents can modulate resistance genes:

MDR → Curcumin, Acetogenin, Haelan

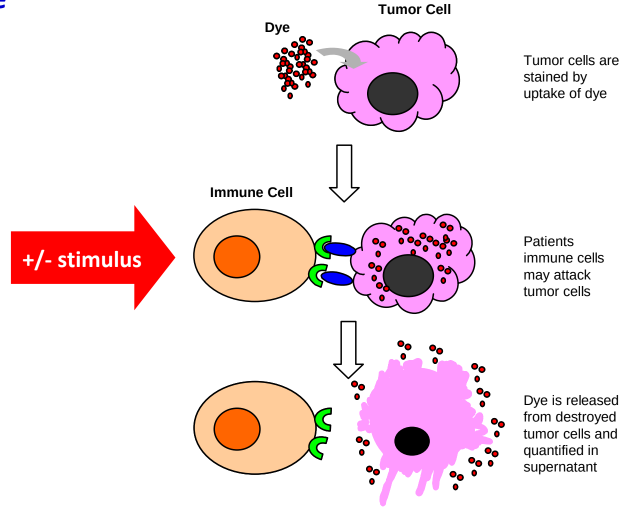
MRP → Artemisinin, Haelan

**GST** → Ellagic acid, Curcumin



# Immune function testing by Cellular NK-Test

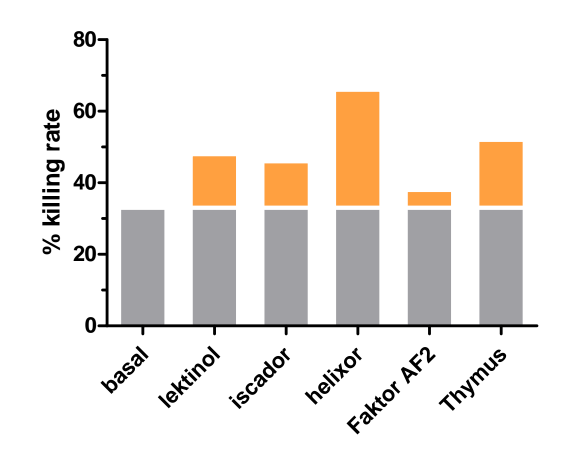
#### **Assay principle**



#### **Cellular NK-test**



#### **Testing of immune stimulative agents**



w/o stimulationafter stimulation

Agent	Included in the assay
Thymus-extract	routinely
Factor AF2	routinely
Iscador mistletoe-extract	routinely
Lektinol mistletoe-extract	routinely
Helixor mistletoe-extract	upon request
Eurixor mistletoe-extract	upon request
Fraxini mistletoe-extract	upon request
Carnivora	upon request
Interleukin 2	upon request



# **Testing for alternative agents**

# Testing is possible for alternative agents with known genetic basis of action:

Agent	
Quercetin	Artemsinin derivatives
IP6 (Inositol-6-Phosphate)	Amygdalin B17
C-statin	Vitamin C
Dammarane sapogenins	Indol-3-carbinol (I3C)
Acetogenin, Graviola	Taurolidine
Haelan951	Ellagic Acid
Curcumin	Arglabin, Laetrile



# Measuring expression of Rhodanese and COX2 expression in tumor cells aids in selection of Amygdalin therapy

# Testing for alternative agents e.g. Amygdalin B17

#### 1. Suppressing expression of COX2 (Cyclooxigenase 2)

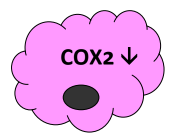
#### COX2:

- Inflammation
- Tumor promotion



CANCER CELL high levels of COX2

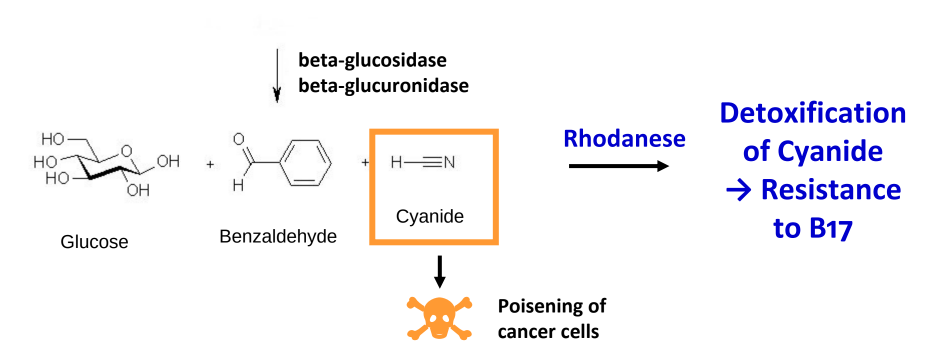
**Amygdalin B17** 



Amygdalin B17 suppresses COX2 expression

# Testing for alternative agents e.g. Amygdalin B17

#### 2. Detoxification by Rhodanese







#### Overexpression of rhodanese in CTC is obviously rare

Rhodanese level	Observed in % of CTCs (n=45)
underexpressed	43 %
base-level (equal to normal cells)	51 %
overexpressed	6 %