

Diagnosis and Treatment of Patients with early and advanced Breast Cancer

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Pathology

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- **Versions 2004–2024:**
**Blohmer / Costa / Fehm / Friedrichs / Harbeck / Huober /
Kreipe / Lück / Maass / Schneeweiss / Sinn / Thomssen / Schmidt**
- **Version 2025:**
Bauerfeind / Kreipe / Sinn

Preamalytics: Fixation

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- **Minimize time to fixation (cold ischemia time)**
- **Minimal fixation time of 6 hours for optimal antigen preservation**
- **Optimal fixation time 6-72 h for core biopsies**
- **Optimal fixation time for resection specimens: 12-72 h**
- **Use of neutral buffered formalin**

	Oxford		
	LoE	GR	AGO
Minimize time to fixation (cold ischemia time)	5	D	++
Minimal fixation time of 6 hours for optimal antigen preservation	5	D	++
Optimal fixation time 6-72 h for core biopsies	5	D	++
Optimal fixation time for resection specimens: 12-72 h	5	D	++
Use of neutral buffered formalin	5	D	++

Use of Breast Cytology

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- **Nipple secretion**
- **Tumor***
- **Cyst**
- **Lymph node***

Oxford		
LoE	GR	AGO
5	D	+
5	D	-
5	D	+/-
5	D	+/-

* Ultrasound-guided core biopsy recommended

Workup: Core Needle Biopsies (US-guided or stereotactic)

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	Oxford		
	LoE	GR	AGO
<ul style="list-style-type: none"> Routine workup in step sections (14G: 1–3 step sections / 11G, 8G: 6–8 step sections) 	5	D	++
<ul style="list-style-type: none"> Correlation with imaging (density, calcifications), use of B-classification 	1b	B	++
<ul style="list-style-type: none"> Frozen section diagnosis on core biopsies 	5	D	--
<ul style="list-style-type: none"> Routine evaluation of ER/PR and HER2 status 	3b	C	++
<ul style="list-style-type: none"> Turn-around time < 24 h (histology) 	5	D	+

Workup: Breast-Conserving Specimens

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- **Slicing perpendicular to the longitudinal axis (or perpendicular to the nipple-peripheral axis in case of spherical specimens)**
- **Systematic sampling, at least 1 tissue block every 1 cm**
- **Inking of resection margins. Sampling of resection margins**
- **Documentation after slicing using specimen radiography, photo documentation or diagram**

Oxford		
LoE	GR	AGO
5	D	++
5	D	++
5	D	++
5	D	+

Workup: Mastectomy Specimens

Oxford

LoE	GR	AGO
-----	----	-----

5	D	++
---	---	----

- | | | | |
|---|---|---|----|
| <ul style="list-style-type: none"> ■ Margins always to be sampled <ul style="list-style-type: none"> ■ Skin close to tumor ■ Deep margin ■ Other margins, if close (< 1 cm) | 5 | D | ++ |
|---|---|---|----|
- | | | | |
|---|---|---|----|
| <ul style="list-style-type: none"> ■ Attention to soft tissue margins in skin sparing mastectomy | 5 | D | ++ |
|---|---|---|----|
- | | | | |
|---|---|---|----|
| <ul style="list-style-type: none"> ■ Routine sampling of uninvolved quadrants, skin above tumor, and retroareolar region | 5 | D | ++ |
|---|---|---|----|
- | | | | |
|--|---|---|----|
| <ul style="list-style-type: none"> ■ Systematic sampling in prophylactic mastectomies (patients with BRCA-1/2 mutation) | 5 | D | ++ |
|--|---|---|----|

Workup: Sentinel Node Biopsy

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	Oxford		
	LoE	GR	AGO
▪ Full workup using step sections of $\leq 500 \mu\text{m}$ on paraffin embedded tissue	5	D	++
▪ Cytokeratin immunohistochemistry			
▪ If suspicious, to detect micrometastases	2b	B	+
▪ For micrometastasis detection after NACT	2b	B	+
▪ As a routine procedure	5	D	+/-
▪ Frozen section (compromises paraffin histomorphology)			
▪ If clinical consequences	5	D	+
▪ If no clinical consequences from frozen section (e.g. cT1 or cT2 and cN0 and BCT)	5	D	-
▪ Imprint cytology instead of, or in addition to frozen section	3b	C	+/-
▪ RT-PCR for epithelial genes	4	D	-
▪ OSNA	3b	B	-

Workup: Intraoperative Pathological Evaluation and Frozen Sections



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	Oxford		
	LoE	GR	AGO
<ul style="list-style-type: none"> ■ Sentinel node biopsy for invasive cancer (compromises final paraffin histomorphology) <ul style="list-style-type: none"> ■ If clinical consequences ■ No clinical consequences ■ Closest margin of resection <ul style="list-style-type: none"> ■ If macroscopically < 1 cm ■ If macroscopically > 1 cm ■ Lesions ≥ 1 cm, without core biopsy ■ Non-palpable lesions or lesions < 1 cm ■ Conservation of fresh tissue (tumor banking) 	5	D	+
	5	D	-
	5	D	+
	5	D	-
	5	D	+
	5	D	--
	5	D	+

Reporting: Histologic Tumor Type

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- **Histologic tumor typing according to WHO-Classification, (5th ed., 2019)**
 - **Partial special differentiation:**
> 50% NST component
and < 50% special tumor type (minor component)
 - **Mixed differentiation:**
> 50% special tumor type
and < 50% NST component
Example: mucinous breast cancer, mixed type
 - **Pure types:**
> 90% special tumor type
Examples: tubular or cribriform Ca.

Oxford

LoE	GR	AGO
3b	C	++



Ductal TNBC: Comparable Survival Rates and Similar Response Rates to Chemotherapy for ER = 0% Compared to ER 1% - < 10%

Reference	Patients	Results
Villegas, S. L. <i>Eur J Cancer</i> 148 , 159–170 (2021) DOI: 10.1016/j.ejca.2021.02.020	Neoadjuvant clinical trial cohorts (n = 2765) comparing neg. ER/PR (< 1%) vs. ER/PR low pos. (ER and/or PR < 9%) vs. strong-pos. (ER or PR ≥ 10%) HR expression.	Low HR-positive, HER2-negative tumours had a similar clinical behavior compared to TNBC, showing high pCR rates and poor survival and also a basal-like gene expression signature. Patients with low HR-positive tumours should be regarded as candidates for therapy strategies targeting TNBC.
Dieci, M. V. et al. <i>Npj Breast Cancer</i> 7 , 101 (2021) DOI: 10.1038/s41523-021-00308-7	406 patients with ER < 10% HER2-negative BC. Pat. Were categorized in ER-negative (ER < 1%; n = 364) and ER-low positive (1–9%, n = 42).	No difference was observed in overall survival (OS) according to ER expression levels (5-years OAS 82.3% vs. 76.7% for ER-negative and ER-low positive BC, respectively, p = 0.8). Our results suggest the use of a 10% cut-off, rather than <1%, to define triple-negative BC (TNBC).
Reddy, S. M. <i>et al.</i> <i>British Journal of Cancer</i> 118 , 17–23 (2018) DOI: 10.1038/bjc.2017.379	Stage I-III TNBC pat. (n = 873) who were disease free at 5 years from diagnosis. Recurrence-free interval (RFI), r.f. survival (RFS), and distant r.f. survival (DRFS) rates were calculated.	After a disease-free interval of 5 years, patients with low hormone receptor-pos. cancers had a higher risk of late events as measured by RFS, and similar risk by RFI or DRFS, compared to TNBC survivors.

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Rare Histological TNBC Subtypes show Divergent Tumor Differentiation Patterns and Clinical Behavior

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Apocrine TNBC

- Luminal phenotype (no basal markers)
- High expression of the androgen receptor
- Low tumor proliferation
- Poor response to chemotherapy
- Better prognosis than ductal TNBC

Metaplastic TNBC

- See chapter 15 Special Situations

Rare and salivary-type TNBC

- Tumors with divergent clinical behavior and specific genetic alterations
- Mostly low tumor proliferation
- Poor response to conventional chemotherapy
- Experimental treatment according to the molecular pathology (e.g. NTREK for secretory ca.)

Apocrine TNBC: More Favorable Survival and Poor Response to Adjuvant Chemotherapy

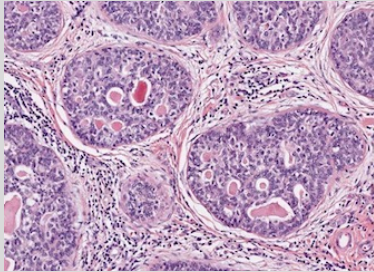
Reference	Patients	Results
<p>Saridakis, A. <i>et al.</i> <i>Ann Surg Oncol</i> 28, 5610–5616 (2021). DOI: 10.1245/s10434-021-10518-9</p>	<p>Women with invasive apocrine cancer were retrospectively identified from the Surveillance, Epidemiology, and End Results (SEER) database. n = 533 triple-negative apocrine cancers were identified.</p>	<p>Half of apocrine tumors are triple negative, but these have more favorable features and much better survival than non-apocrine triple-negative cancers. Compared with non-apocrine triple-negative, apocrine triple-negative patients were much older, with smaller, lower-grade tumors and much better survival (86% vs. 74%).</p>
<p>Montagna, E. <i>et al.</i> <i>Breast</i> 53, 138–142 (2020). DOI: 10.1038/s41523-021-00308-7</p>	<p>406 patients with ER < 10% HER2-negative BC. Pat. Were categorized in ER-negative (ER < 1%; n = 364) and ER-low positive (1–9%, n = 42).</p>	<p>The outcome of selected apocrine triple negative breast cancer patients who did not received adjuvant chemotherapy is excellent and supports a treatment de-escalation.</p>
<p>Mills, A. M., <i>et al.</i> <i>Am J Surg Pathol</i> 40, 1109–1116 (2016). DOI: 10.1097/pas.0000000000000671</p>	<p>All pure apocrine carcinomas diagnosed during a 10-year period were reviewed, and clinicopathologic characteristics were compared with a control group of 26 non-apocrine TNBC cases. Twenty apocrine carcinomas were identified (~ 0.8% of all breast cancers).</p>	<p>Apocrine TNBC had a favorable clinical prognosis, with 80% of patients showing no evidence of disease-related morbidity or mortality (mean follow-up: 45.2 mo). Pure apocrine carcinomas represent a clinicopathologically distinct subgroup of triple-negative breast cancer characterized by AR positivity.</p>

Rare and Salivary-type TNBC: Tumors with Divergent Clinical Behavior and Specific Genetic Alterations

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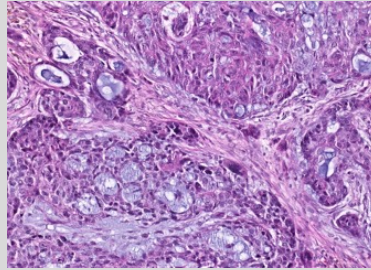
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Adenoid-cystic carcinoma



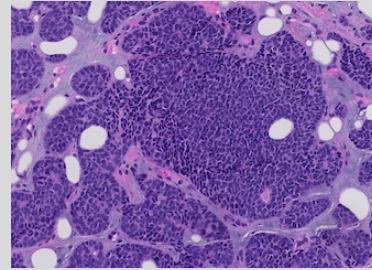
MYB-NFIB
MYBL1 rearrangements
MYB gene amplification

Secretory carcinoma



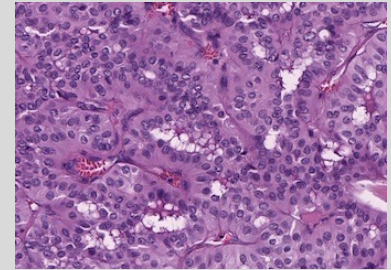
ETV6-NTRK3
gene fusions

Polymorphous carcinoma



PRKD1 E710D
PRKD1/PRKOZ/PRKD3
rearrangements

Tall cell carcinoma with reversed polarity



IDH2 hotspot mutations

Reporting: Grade of Malignancy

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	Oxford		
	LoE	GR	AGO
<ul style="list-style-type: none"> Use of Nottingham grading system (Elston & Ellis 1991) for all types of invasive breast cancer (incl. status post neoadjuvant systemic therapy) 	5	D	++
<ul style="list-style-type: none"> In case of very little tumor tissue, pure nuclear grading or additional criteria, such as Ki-67 proliferation fraction, may be used 	5	D	++
<ul style="list-style-type: none"> Grading of DCIS, e.g. according to WHO-Classification, (5th ed., 2019) 	5	D	++
<ul style="list-style-type: none"> Reporting of tumor grade in numeric form (e.g. G3) 	5	D	++

Reporting: Tumor Size and Total Extent of Tumor

Oxford

LoE	GR	AGO
-----	----	-----

5	D	++
---	---	----

- Reporting of invasive tumor size taking into account macroscopic and histologic findings and clinical imaging results

5	D	++
---	---	----

- Additional reporting of total extent of invasive carcinoma in case of satellite nodules or multifocality

5	D	++
---	---	----

- Reporting of size of non-invasive component (DCIS or LCIS) when DCIS or LCIS component is extensive (more than 2x invasive Ca)

Reporting: pTNM

Oxford

LoE	GR	AGO
5	D	++

- Use of current UICC classification (8th ed.)

pT 1-3: Invasive tumor size (largest focus in case of multifocality or multicentricity)

pT4: Invasion of dermis alone does not qualify as pT4. Criteria for pT4a/b/c/d must be met.

pT4d: Negative skin biopsy does not rule out pT4d (inflammatory carcinoma).

pM: pM1 indicates any non-regional disease, except 2nd primary contralateral.
Use of MX is not recommended.

Reporting: Margins of Resection and R-Classification

Oxford

LoE	GR	AGO
-----	----	-----

5	D	++
---	---	----

- Evaluation of distance to all resection margins macroscopically and close margins histologically (< 1 cm)

5	D	++
---	---	----

- Reporting of minimal distance to resection margin and its topography

5	D	++
---	---	----

- R-Classification

R0: No residual tumor

R1: Microscopic invasive or noninvasive carcinoma involving resection margin

RX: Presence of residual tumor cannot be assessed (e.g. tumor in multiple specimens)



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Reporting: Lymphovascular Invasion

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- **L1: Lymphovascular invasion**
L0: No lymphovascular invasion
- **IHC for evaluation of lymphovascular invasion**
- **Differentiation of peritumoral and extensive lymphovascular invasion**
- **Reporting of venous invasion (V0/V1) optional, prognostic significance not established**

Oxford		
LoE	GR	AGO
5	D	++
3b	C	-
3b	C	++
5	D	+

Reporting: Evaluation of Tumor-Infiltrating Lymphocytes (TIL)

Oxford

LoE	GR	AGO
5	D	+/-

- **Identification of tumors with predominant lymphocytic infiltrate (> 50%) in tumor stroma (according to Salgado et al.*)**

Consider only lymphocytic infiltrate in tumor stroma and not at the invasion front

Do not consider central fibrosis and necrotic areas

Report average of lymphocytic infiltrate as percentage

- * Salgado, R., Denkert, C., Demaria, S., Sirtaine, N., Klauschen, F., Pruneri, G., et al. (2014). The evaluation of tumor-infiltrating lymphocytes (TILs) in breast cancer: recommendations by an International TILs Working Group 2014. *Annals of Oncology*

Reporting: Evaluation after Neoadjuvant Chemotherapy

Oxford

LoE GR AGO

	LoE	GR	AGO
▪ Identification of tumor bed, otherwise ypTX	4	D	++
▪ Reporting of tumor size as total extent of tumor bed area involved by infiltrates of residual vital invasive carcinoma	4	D	++
▪ pCR when absence of invasive Ca. and absence of angioinvasion or LN metastases. Presence of ypTis should be recorded	2b	D	+
▪ Use of IHC to identify tumor residues (lymphnodes)	2b	B	+/-
▪ Reporting of ypTN after neoadjuvant systemic therapy	5	D	++
▪ Repeat IHC for ER, PR, and HER2	5	D	+/-
▪ Intraoperative frozen section (reduced sensitivity)	5	D	-
▪ Tumorregression-Scores: RCB-Score or Sataloff-Score	4	D	+/-

Predictive Pathology of Endocrine Responsiveness

Oxford		
LoE	GR	AGO
1a	A	++
1b	A	+
1b	A	+

- Immunohistochemical detection of estrogen- and progesterone-receptors in paraffin-embedded tissue; scored as percentage of positive tumor cell nuclei (ER positive if $\geq 1\%$, low positivity $\geq 1\%$ to 10% ; PR positive if $\geq 10\%$)**
- Detection of endocrine responsiveness by Ki-67 decrease to $\leq 10\%$ after 3-4 weeks of preoperative endocrine therapy in primary breast cancer**
- Detection of secondary, i.e. acquired endocrine resistance by analysis of activating ESR-1 mutations in liquid biopsy or metastatic tissue**

HER2-Analysis by IHC

Oxford

LoE	GR	AGO
-----	----	-----

- | | | | |
|--|----|---|----|
| <ul style="list-style-type: none"> 3+ staining pattern: HER2+ if strong complete circular membrane staining of > 10% invasive cells | 1a | A | ++ |
| <ul style="list-style-type: none"> 2+ staining pattern: If > 10% circular but moderate/weak membrane staining or ≤ 10% strong staining, U-shaped staining in micropapillary carcinoma: ISH required (CISH, SISH, FISH) | 1a | A | ++ |
| <ul style="list-style-type: none"> 1+ staining pattern: with > 10 % incomplete membrane staining that is weak or barely perceptible (caveat: reproducibility). | 1a | A | + |
| <ul style="list-style-type: none"> 0 grade staining: to be confirmed by second determination in case that Trastuzumab-Deruxtecan treatment* is considered | 5 | D | ++ |
| <ul style="list-style-type: none"> HER2-low: 1+ oder 2+ /ISH negativ | 1b | A | ++ |

* Due to heterogeneity and therapeutic relevance

HER2-Analysis by ISH when IHC 2+

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Oxford

LoE GR AGO

- **Single-Color In-Situ-Hybridisation (ISH):**
 - HER2+ if signal counts ≥ 6 in at least 20 cohesive cells
 - negative if signal counts < 4 signals/nucleus
 - 2-Color ISH recommended for ≥ 4 and < 6 signals/nucleus

- **Two-Color In-Situ-Hybridisation (ISH):**
 - Group 1: Ratio ≥ 2.0 and signals/nucleus ≥ 4.0 -> HER2+
 - Group 2: Ratio ≥ 2.0 and signals/nucleus < 4.0
-> HER2- (no benefit of anti-HER2 therapy)
 - Group 3: Ratio < 2.0 and signals/nucleus ≥ 6.0
-> HER2+ (but benefit of anti-HER2 therapy not certain)
 - Group 4: Ratio < 2.0 and signals/nucleus ≥ 4.0 und < 6
-> HER2- (no benefit of anti-HER2 therapy)
 - Group 5: Ratio < 2.0 und signals/nucleus < 4.0 -> HER2-

3a C ++

3a D ++

HER2 testing by validated dual-probe ISH assay when IHC = 2+

Batch controls and on-slide controls show appropriate hybridization

HER2/CEP17 ratio ≥ 2.0

HER2/CEP17 ratio < 2.0

Group 1
Average HER2
copy number ≥ 4.0
signals/cell

Group 2
Average HER2
copy number < 4.0
signals/cell

Group 3
Average HER2
copy number ≥ 6.0
signals/cell

Group 4
Average HER2
copy number ≥ 4.0
- < 6.0 signals/cell

Group 5
Average HER2
copy number < 4.0
signals/cell

mostly

mostly

mostly

HER2
positive

HER2
negative

HER2
positive

HER2
negative

HER2
negative

HER2 Testing on Core Biopsies

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False positive immunohistochemical labeling may occur in core biopsies. Therefore, methods of individual laboratories should be validated by comparison of core biopsies and resection specimens. Background staining should be evaluated by comparison with normal duct epithelium.

Alternatively, all G1 and G2 cases with HER2 3+ in core biopsies may be analyzed by ISH or may be re-evaluated in the resection specimen.

False positivity is likely when HER+ was reported in G1 tumors of the following types: Infiltrating ductal or lobular carcinoma, ER and PR positive, Tubular (at least 90% pure), Mucinous (at least 90% pure) Cribriform (at least 90% pure), Adenoid cystic carcinoma (90% pure).

In case of discrepancy between core biopsy and specimen, the HER2 overexpressing sample should be re-evaluated by a different method. If still discrepancy – anti-HER2-treatment if amplified in one of both samples. Expected rate of HER2-overexpression: 15% HER2 positive.

Additional Special Studies: Molecular Analysis of HER2 Status

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<ul style="list-style-type: none"> Therapy decisions should only be based on IHC and ISH 	1a	A	++
<ul style="list-style-type: none"> Evaluation of HER2 using validated gene expression test kits 	3b	B	-
<ul style="list-style-type: none"> Evaluation of HER2-amplification by RNA-sequencing 	5	D	-
<ul style="list-style-type: none"> Use of molecular HER2-testing for subtyping 	3b	B	+/-

Special Studies: Evaluation of Ki-67 Score

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	Oxford		
	LoE	GR	AGO
▪ Counting of tumor nuclei at the invasion front	5	D	++
▪ Semiquantitative eyeballing or counting of labelled cells in core needle biopsies	2	A	++
▪ Consideration of weakly stained tumor nuclei	5	D	++
▪ Reporting of Ki-67 positive nuclei as percentage	5	D	++
▪ Establishing of laboratory standards and cut-off values	5	D	++
▪ Use of image analysis for objective Ki-67 evaluation	5	D	+
▪ Determination of Ki-67 dynamics after short term (2-4 weeks) endocrine therapy*	1b	B	+

* See chapter Neoadjuvant Systemic Therapy

Predictive PD-L1 Determination in Metastatic Triple Negative Breast Cancer

Oxford

Immunohistochemical assay

LoE GR AGO

Metastatic or primary tumor tissue

2 A ++

Detection with antibodies equivalent to registration trials

3 B +

Combined positive score (CPS) for pembrolizumab indication

2 A ++

Divide: $\frac{\text{positive tumor cells} + \text{macrophages} + \text{lymphocytes}}{\text{number of tumor cells}} \times 100$

Cut-off value: CPS \geq 10

1b A

Immune Score (IC) for atezolizumab indication: Cytoplasmic staining of the leucocyte stromal infiltrate (lymphocytes, macrophages, plasma cells, granulocytes outside of abscesses) in relation to the tumor volume

2 A ++

Cut-off value: IC \geq 1%

1b A



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Mutational Studies* in mBC: „Precision Medicine“ for Targeted Therapies

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Altered genes	Therapeutic relevance	Gene region	Material	Oxford		
				LOE	GR	AGO
BRCA1, BRCA2	Olaparib, Talazoparib Olaparib	All exons	Germline: Blood cells	1b	A	++
			Somatic: Tissue	2b	B	+
PALB2	Olaparib		Germline: Blood cells	2b	B	+
PIK3CA	Alpelisib / Inavolisib	Exons 7, 9 and 20	Primary tumor, metastases, plasma	1b	A	++
AKT1, PTEN, PIK3CA	Capivasertib		Primary tumor, metastases, plasma	1b	A	+
HER2-mutation (independent of HER2-status)	Neratinib, lapatinib	Kinase- and extracellular domains; S310, L755, V777, Y772_A775dup	Primary tumor, metastases, plasma particul. lobular BC	4	C	+/-
ESR1	Resistance against AI Response to Elacestrant	Exons 4, 7 and 8	Metastases, plasma	2b	B	+
			Metastases, plasma	1b	B	++
NTRK gene fusion	Larotrectinib, entrectinib	Fusion- and splice variants	Tumor tissue, particul. secretory breast cancer	2a	B	+
MSI	Pembrolizumab	Microsatellite-instability	Tissue	2a	B	+

* Ideally panel diagnostics