

HER2 Signaling Function Test

# Background

HER2 gene (ERBB2) amplification and/or HER2 protein overexpression is detected in approximately 15-20% of breast cancers and associated with more aggressive disease progression, metastasis, and a poorer prognosis [1-4]. The use of HER2-targeting agents, such as monoclonal antibodies, trastuzumab (Herceptin) and pertuzumab (Perjeta), significantly improves clinical outcomes in HER2+ patients [4, 5]. Currently, a patient's eligibility for HER2-targeted therapies is determined by their IHC- or FISH-based HER2 testing scores [4]. However, clinical trials have indicated a weak correlation between HER2 expression levels and HER2 targeted therapy benefit [6, 7]. Other biological factors, such as HER2 signaling activity, may be important to measure, in addition to expression and amplification of HER2, when identifying patients eligible for HER2 therapies.

A new functional cellular analysis platform, the CELx HER2 Signaling Function (CELx HSF) Test, which uses a label-free impedance biosensor to measure HER2 signaling activity in live tumor cells, has been developed by Celcuity [8, 9] (Figure 1). This test has been applied to tumor cell samples from 89 patients' fresh HER2- breast cancer tissue (DAKO 0 or 1+) and found abnormal HER2 signaling activity in 16 of 89 (18%; 95% CI=11%-27%) patients. This work that has been published recently [9], which suggests a new sub-group of HER2- breast cancer patients may benefit from anti-HER2 therapy.

In order to extend the understanding of any benefit this new sub-group may receive from HER2 therapy, the current study set out to quantify the percentage of HER2-driven signaling activity two HER2 mAb's, trastuzumab (Tz) and pertuzumab (Pz), both alone and in combination, could inhibit two groups of live cells ex vivo: 1) HER2- breast tumor cells with abnormal HER2-driven signaling (n=5); and 2) HER2+ breast cancer cell lines (n=4).

# Methods

Specimens and Cell Culture: Epithelial cells derived from five HER2- primary tumors (IHC 0 or 1+) and HER2+ breast cancer cell lines, each with comparable levels of abnormal HER2-driven signaling, were used in this study. Methods for tissue extraction and primary cell culture are essentially as described previously [10, 11]. Modifications can be found in our recent publication [9]. All cell lines were maintained according to ATCC recommendations and authenticated by ATCC in March 2016.

Flow Cytometry: Flow cytometry was performed on a BD FACSCalibur™ system using cells harvested at the time of the CELx HSP test using FACS markers listed in Table 2. Flow cytometry results are 100% concordant to the standard clinical IHC test evaluations for HER2, ER, and PR markers that were provided for each specimen by the clinic.

CELx HSF Test and Data Analysis: Real time live cell response to specific HER2 agonists (NRG1b or EGF) and with or without an antagonist (trastuzumab, pertuzumab, or the combination of the two antibodies) was measured and quantified using an xCELLigence RTCA impedance biosensor (ACEA Biosciences). Clinically relevant concentrations of agonist and each HER2 mAb were used. From these responses, the net amount of HER2 participation in HER2 signaling initiated by the HER2 agonists ("HER2s") was determined (Figure 1). To classify a HER2- sample as having abnormal signaling, a test cut-off value of 250 was established [8, 9]. The percentage of inhibition of NRG1-driven HER2 CELx signal by trastuzumab alone, pertuzumab alone, or their combination in each sample was calculated and compared between the HER2- primary cell group (n=5) and the HER2+ cell line group (n=4).

#### Table 1. Antibodies Used in Flow Cytometry

Description of Antibody	Vendor
Mouse anti-human HER2-phycoerythrin (PE), clone 24D2	Biolegend, San Diego, CA
Mouse anti-human HER1 conjugated with AlexaFluor 647, clone EGFR.1	BD Biosciences, San Jose, CA
Mouse anti-human HER3 conjugated with PerCP-sFluor710, clone SGP1	eBioscience, San Diego, CA
Mouse anti-human EPCAM conjugated with AlexaFluor 488, clone MH99	eBioscience, San Diego, CA
Mouse anti-human Claudin4 conjugated to PE, clone 382321	R&D Systems, Minneapolis, MN
Rat anti-human CD49f conjugated to PerCP/eFluor710, clone eBioGoH3	eBioscience, San Diego, CA
Mouse anti-human CD10 conjugated to Allophycocyanin (AP), cloneHL10a	BioLegend, San Diego, CA
Rabbit polyclonal anti-human estrogen receptor alpha (ER $lpha$ ) conjugated to AlexaFluor488	Bioss, Woburn, MA
Mouse anti-human progesterone receptor (PRG) conjugated to eFluor660, clone KMC912	eBioscience, San Diego, CA

\*All epitopes were extracellular with the exceptions of ER and PR. All antibodies were purchased from companies as listed who provided empirical demonstration of each of the antibodies for our applications

#### References

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# Inhibition of abnormal HER2-driven signaling by two HER2 targeted antibody drugs tested ex vivo in live primary HER2- breast cancer cell samples and HER2+ cell lines

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### Table 2. Examples of HER2-Negative (HER2<sup>-</sup>) Patient Samples with Abnormally High HER2-Dependent Signaling (HER2<sub>s</sub>+) Identified by CELx HSF Test

HER2 <sup>-</sup> Patient ID	NRG1-driven HER2-dependent CELx Signal Units	EGF-driven HER2-dependent CELx Signal Units	Total HER2-dependent CELx Signal Units	CELx Test Result
R39 (tumor)	475	88	563	Abnormal
R20 (tumor)	409	120	529	Abnormal
R160 (tumor)	332	99	430	Abnormal
R82 (tumor)	272	40	312	Abnormal
R95 (tumor)	227	44	271	Abnormal
R25 (tumor)	238	29	267	Abnormal
R71 (tumor)	228	23	252	Abnormal
R22 (tumor)	1	78	79	Normal
R62 (healthy)	31	7	38	Normal
SKBr3 (HER2+ cell line)	401	143	544	Abnormal

In one of our studies, CELx HSF tests were performed on 34 tumor cell samples from patients with breast cancer classified as HER2<sup>-</sup> to measure HER2 pathway stimulation and signal specificity [9]. For comparative purposes, 16 primary breast epithelial cell samples from healthy patients, and DAKO IHC test standard breast cancer cell lines SKBr3 (IHC 3+) and MDA-231 (IHC 0) were also subjected to CELx HSF Tests. NRG1b-induced and EGF-induced CELx signals for tumor and healthy primary cells and a HER2<sup>+</sup> reference cell line (SKBr3) (DAKO test score 3+) are summarized in the table.

In this study, the CELx HSF test identified 7 of 34 (~20%) HER2- patient samples having abnormally high HER2 signaling activities.

#### Figure 3. CELx Test Identifies Four Different Subgroups for HER2 Signaling

Examples of CELx HSF test results (HER2 $\pm$ /HER2<sub>s</sub> $\pm$ ): use of NRG1 (ligand) and pertuzumab (antagonist) to ID net HER2-driven signaling activity. Within HSF+ subgroups differential response to drug is demonstrated.

HER2+/S- CL (BT474)	HER2-/S+ (Patient R39)	HER2-/S- (Patient R49)
800	800	800
600	600	600
400	400	400
200	200	200
O NRG1 NRG1+P Inhibition (Δ)	0 NRG1 NRG1+P Inhibition (Δ)	0 NRG1 NRG1+P Inhibition (Δ)
d widely in the HER2+ of highest HER2+ cell lines	<ul> <li>HER2- primary tumor samp similar range of NRG1 initia cell lines</li> </ul>	oles, R39 and R49, show a ated activity as the HER2+
	HER2+/S- CL (BT474) 800 600 400 200 0 NRG1 NRG1+P Inhibition (A) MRG1 HER2+ of highest HER2+ cell lines	HER2+/S- CL (BT474) 800 600 400 200 0

- Difference between NRG1 and NRG1 + P above is amount of signaling involving HER2 in HER2/HER3 dimerization
- Pertuzumab inhibits 50% of the SKBR3 signal
- None of BT474's limited activity is inhibited
- The HER2-driven activity ( $\Delta$ ) for R39 is hearly identical
- to SKBR3's High Pz inhibition is similar to other HER2/HER2<sub>s</sub>+
- R49's low signaling is typical of 80% of HER2- patient tumors

#### Table 3. Comparison of Inhibition of NRG1-driven HER2 CELx Signals by Trastuzumab (Tz) or Alone or in Combination

Average % Inhibition of NRGI-driven HER2 Signaling Activity				
HER2 mAb	HER2+/HER2 <sub>s</sub> + Cell Lines (n=3)	HER2-/HER2 <sub>s</sub> + Primaries (n=5)		
Trastuzumab	19%	44%		
Pertuzumab	62%	73%		
Trastuzumab + Pertuzumab	87%	81%		

Pertuzumab and trastuzumab alone were each more effective in the HER2- group than in HER2+ group.



#### Figure 4. Examples of Comparison of NRG1-driven HER2 Signaling Inhibition by Trastuzumab (Tz) or Pertuzumab (Pz) Alone or in Combination





# Summary of Results

- All cell samples tested with pertuzumab and trastuzumab had comparable, and abnormal, levels of NRG1 activated HER2-driven signaling.
- In each sample, the two mAbs (trastuzumab and pertuzumab) inhibited a higher percentage of signaling in combination than either mAb alone; no interference effects between the two mAbs were detected.
- Pertuzumab and trastuzumab alone were each more effective in the HER2- cell samples than in the HER2+ ones.

# Conclusions

- These findings provide strong evidence that the two HER2 mAbs (trastuzumab and pertuzumab) used to treat HER2+ breast cancer patients are as effective in blocking abnormal HER2-driven signaling function *ex vivo* in HER2- primary cells with abnormal signaling as they are in HER2+ cells with abnormal signaling.
- Overexpressed HER2 receptor may thus not be a required condition for breast cancer patient response to HER2 mAbs.
- These results suggest a new sub-group of breast cancer patients, HER2- with abnormal HER2 signaling (HER2-/HER2<sub>s</sub>+), identified with the CELx HER2 Signaling Function (HSF) Test, may benefit from the addition of HER2 mAbs to current therapeutic regimens.
- Additional studies to confirm these findings are currently underway.