

Evaluation of pan-HER and c-Met inhibitors tested ex vivo in live primary HER2- breast cancer cells with hyperactive c-Met and (HER) ErbB family signaling

Laing LG, Burns DJ, Khan SE, Kharbush S, Soltani SM, MacNeil IA, Rich BE, Brass K, Broege AM, Mutka S, Sullivan B – Celcuity Inc., 16305 36th Ave N, Suite 100, Minneapolis, MN 55446, USA

Background

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.

HER2 gene (ERBB2) amplification and/or HER2 protein overexpression is detected in approximately 15–20% of breast cancers and is associated with more aggressive disease progression, metastasis, and a poorer prognosis.¹⁻⁴ Agents targeting HER2, such as trastuzumab, lapatinib, and pertuzumab, significantly improve clinical outcomes in HER2⁺ patients.^{4,5} Currently, a patient's eligibility for HER2 targeted therapies is determined using IHC or FISH HER2 tests.⁴ However, clinical trials have indicated a weak correlation between HER2 expression or amplification levels and HER2 targeted therapy benefit.^{6,7}

c-MET is the cognate receptor for Hepatocyte Growth Factor (HGF). MET amplification and HGF overexpression have emerged as mechanisms by which cancers become resistant to HER family therapies.^{8,9} c-Met and HER family signaling pathways also participate in extensive cross-talk that can drive cancer progression.¹⁰ Phase II trials with c-Met targeted therapies, either cabozantinib or onartuzumab, in combination with erlotinib, showed promising improvement in PFS.¹¹ However, Phase III clinical trials studying c-Met targeted therapies using MET overexpression as an enrollment criteria and studies in which patients were randomly selected to receive combination c-Met/HER1 treatment failed to demonstrate clinical efficacy.¹¹ Because c-Met expression level using IHC as a clinical pathology diagnostic marker fails to identify a population responsive to c-Met targeted therapies, an alternative approach is required to identify patients with dysfunctional c-Met signaling who will respond to these therapies.

We previously developed a test to identify patients with abnormal HER2 signaling.^{12,13} To elucidate the role of c-Met signaling and its involvement with HER family signaling as a cancer driver, a new assay using an impedance biosensor and live cells derived from each patient's tumor, the CELx multi-pathway signaling function (CELx MP) test, was developed.

The CELx MP test measures ex vivo real-time live cell response to specific HER family and c-Met agonists to diagnose breast tumors with hyperactive HER1, HER2, HER3, HER4, and c-Met signaling activity. A recent study quantified c-Met and HER-driven signaling activity in epithelial cell samples derived from fresh breast tumor specimens obtained from 79 HER2-negative breast cancer patients. Of the cell samples tested, 19 of 79, (24.1%; 95% CI=16–35%) had both hyperactive c-Met signaling and at least one hyperactive signaling HER-family receptor.

Using primary breast cancer cells with hyperactive c-Met and HER family signaling and the CELx MP test, the current study set out to:

- 1. Determine the IC50 values of six pan-HER inhibitors and five c-Met inhibitors.
- 2. Characterize the efficacy of combinations of each pan-HER inhibitor with each c-Met inhibitor.

Methods

Specimens: Epithelial cells from six HER2-negative tumor specimens with hyperactive c-Met and HER-driven signaling were obtained.

Cell Culture: Methods for tissue extraction and primary cell culture* are essentially as described previously.^{14,15}

Flow Cytometry: Flow cytometry of all cell samples was performed on a BD FACSCalibur using cells harvested at the time of the CELx MP test. Flow cytometry results are 100% concordant to the standard clinical IHC test evaluations for HER2 that were provided for each specimen by the clinic.

CELx MP Test: Real-time live cell responses to specific HER3 and HER1 agonists (NRG1b and EGF) or c-Met agonist (HGF) with and without antagonists were measured and quantified using an xCELLigence RTCA impedance biosensor (ACEA Biosciences). From these responses, the net amount of HER2 participation in HER-family signaling initiated by HER receptor agonists was determined.^{13,14} Samples with HER2 signaling activity levels above a previously determined cut-off value of 250 signaling units that was attenuatable with a HER2 dimer blocker were identified as abnormal. Samples with a c-Met signal greater than 250 were characterized as hyperactive.

Determining IC50 values: A 1,000-fold, 5-point dose response of a panel of antagonists for HER-family and c-Met was run against the corresponding agonist stimulus (NRG1b/EGF for HER-family activation and HGF for c-Met activation) in a live-cell CELx assay. Response as a percent of maximum response of the agonist without an antagonist was plotted in GraphPad Prism software and IC50 values were derived using nonlinear regression. IC50 values were determined in six primary breast cancer cells and the values were averaged.

Figure 1. Platform Biosensor Sensitivity Enables Quantification of HER and c-Met Signaling Real-Time in Live Cells



* De-identified excess surgically resected human breast cancer tissue was received from multiple clinical sites located across the United States. Liberty IRB (Columbia, MD) determined that this research did not involve human subjects as defined under 45 CFR 46.102(f) and granted an IRB exemption. Liberty IRB has full accreditation with the Association for the Accreditation of Human Research Protection Programs (AAHRPP)

Figure 2. HER2⁻ Abnormal Signaling by CELx HSF Test

Representative CELx Time-Course Curves



CELx time-course curves representing a high, abnormal HER2 signal in a high responder (R39) and a low HER2 signal in a non-responder (R58). In this display, curves of NRG1 stimulation in the absence versus presence of HER2 dimer blocker (10µg/mL) are presented. The data show that the high NRG1b responder has more than 10 times greater signal than the low responder, indicating the test has a large dynamic range.





(A) Comparison of expression levels of HER2, HER3 determined by flow cytometry with CELx test score for 79 patient primary breast cancer cells. Fluorescence values for HER2 DAKO 3+ cells is approximately 2,000 MFC units.

B) Percent response of stimulation of NRG1b/EGF signal (two-growth factor cocktail) when treated with a 5-point, 1,000-fold titration of the six indicated pan-HER antagonists. (Table 1, right) Average IC₅₀ (nM) of pan-HER inhibitors from six primary breast cancer cell samples found with hyperactive HER-family and c-Met signaling. IC50 values for pan-HER antagonists against the NRG1b/EGF stimulation determined from real-time live cell assay align with what is reported in cellfree assays.

Figure 4. Determining IC50 Values of c-Met Inhibitors Against HGF Agonist in a Real-Time Live Cell Assay



(A) Comparison of expression levels of HGF receptor (c-Met) determined by flow cytometry with HGF test score from a set of primary breast cancer cells from 79 patients

(B) Percent response of stimulation of HGF signal when treated with a 5-point, 1,000-fold dose response of the five indicated c-Met antagonists. (Table 2, right) Average IC₅₀ (nM) of c-Met inhibitors from six primary breast cancer cells with hyperactive ErbB-family and c-Met signaling.

IC50s for c-Met antagonists against the HGF response determined from real-time live cell assay align with what is reported in cell-free assays.

Results

Pan-HER Inhibitors	IC ₅₀ (nM)
Poziotinib	1.23
Neratinib	4.81
Ibrutinib	13.10
Dacomitinib	22.06
Sapitinib	41.28
Lapatinib	137.27

*Poziotinib average IC50 derived from dose response on three primary breast cancer cell samples.



c-Met Inhibitors	IC ₅₀ (nM)
Capmatinib	3.10
Savolitinib	3.56
Tepotinib	14.70
Cabozantinib	27.36
Crizotinib	28.21

Figure 5. Combinations of Pan-HER and c-Met Inhibitors Effectively Block NRG1b/EGF/HGF Agonist Cocktail in Real-Time Live Cell CELx MP Test



Pan-HER Inhibitors	Average Inhibition (%) with Different c-Meti's	c-Met Inhibito	ors Average Inhibition (%) with Different HE
Poziotinib	100	Capmatinib	94
Neratinib	100	Savolitinib	98
Ibrutinib	99	Tepotinib	96
Dacomitinib	100	Cabozantini	b 99
Sapitinib	98	Crizotinib	100
Lapatinib	80		

(A) CELx MP Test data showing activation by a NRG1b/EGF/HGF cocktail (N/E/H, red line) with c-Met (tepotinib) or pan-HER (neratinib) inhibitors alone or in combination. Antagonists were added 6 hours after initial seeding followed by an incubation period prior to growth factor addition. (B) (Table 1, left) Average percent inhibition of the N/E/H signal with five different c-Met inhibitors against each pan-HER inhibitor. (Table 2, right) Average percent inhibition of the N/E/H signal with six different pan-HER inhibitors against each c-Met inhibitor. Targeting c-Met and HER signaling pathways effectively eliminates signaling in the CELx signaling function assay.

Summary of Results

- 1.23 nM 137.27 nM.
- The average IC50 values for the individual c-Met inhibitors ranged from 3.10 nM – 28 nM.
- In drug efficacy studies, an average of at least 80% of the ErbB and c-MET signaling activated by NRG1b, EGF, and HGF co-stimulation was inhibited by each combination of c-Met and pan-HER inhibitors.

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Results

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			$C 2 G \Lambda \perp N / \Box / \Box *$
			CZO4 + IN/E/II
and the second se			C264 + N/E/H + tepotinib
			C264 + N/E/H + neratinib
			C264 + N/E/H + neratinib + tepotinib
	:	:	
	1	2	3 4
		Time (ho	urs)

The average IC50 values for the individual pan-HER inhibitors ranged from

Conclusions

- The CELx MP test using live cells measures IC50 values comparable to those derived using cell-free methods.
- Every combination of pan-HER and c-Met inhibitors provided comparably high (at least 80%) levels of inhibitory activity effect *ex vivo*.
- These results suggest the sub-group of HER2-negative breast cancer patients diagnosed with coincident hyperactive c-Met and ErbB signaling by the CELx MP test may respond to virtually any pan-HER and c-MET inhibitor combination.
- Studying combinations designed to minimize drug toxicities without sacrificing efficacy should thus be possible.
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