Only trace amounts of 1:1 and 2:1 complexes were detected for zani or the order complexes with some differences of molecular weight distribution.

Zanidatamab (zani) simultaneously binds two distinct epitopes of the oncogenic cell surface receptor HER2. Zani is currently being evaluated in global Phase 1, Phase 2 and registration-enabling clinical trials as a potential activity over tras or combination of tras and pert. We present results from a structural characteristics of zani:HER2 complexes.

The extracellular domain of the HER2 receptor comprises four domains with receptor clustering, removal of HER2 from the cell surface, and potent effector function. Preclinical data shows that zani can exhibit differentiated structural and imaging studies which showcase the differentiated structural characteristics of zani:HER2 complexes.

**Zanidatamab: A Biparatopic Anti-HER2 Antibody**

**Zanidatamab:HER2 Do Not Form Simple 1:1 Paired Complexes**

Complexes of zani with HER2 appeared as strings of molecular assemblies on the Cryo-electron micrographs. A model of HER2 (cyan) bound to zani was determined to approximate a 8 Å resolution from a dataset of about 25,000 particles. Only the Fab (blue ribbons) bound to domain 2 of HER2 and scFv (green ribbons) bound to domain 4 of HER2 are resolved. The relative orientation of the Fab and scFv does not sterically allow for a 1:1 pairing of HER2 and zani. Instead, 2 copies of zani are expected to bind HER2 as shown in the example 2D class average and 3D model, with the rest of the structure appearing as a diffuse halo. The unresolved domains of two bound zani molecules are presented in the 3D structure as cartoons.

**Zanidatamab Induces Distinctly Large Clusters of HER2 On Target Cell Surface**

In a 3D confocal spinning microscope imaging study with breast cancer cell line SKBR3 that over-expresses HER2, fraction of cells treated with zani showed strikingly large HER2 ‘caps’ that are polarized to one side of the cell membrane. While all α-HER2 antibodies tested induced receptor aggregation and microcluster formation, such receptor caps were largely absent in cells treated with tras, pert or their combination (See Abstract number 1005).

**Computational Modeling Helps Understand Characteristics Of Zani:HER2 Complexation**

Hexamerization of IgG fc promotes C1q engagement and CDC activity. In light of the CDC activity observed for zani (See Poster #1005), we evaluated if a HER2:zani complex could form a six-fold cyclic structure. Applying symmetry operations to a model of HER2 bound to two copies of zani (top left) based on the structure of HER2 + tras/pert complexes (PDB ID: 1N8Z, 1S78, 6OGE) resulted in a six-fold ring structure (bottom left) with HER2 naturally aligned in a plane, akin to presentation on the cell membrane (below). The fc is aligned in a configuration suitable for interaction with C1q (PDB ID: 6J2C). Symmetry limitations did not allow an equivalent model of HER2 with the combination of tras and pert.

**Zanidatamab Induced Clusters Are Larger Than Those Formed By Trastuzumab (T), Pertuzumab (P) Or T+P**

We used single molecule localization microscopy (SMLM) with a real-time drift-stabilized direct Stochastic Optical Reconstruction Microscope (dSTORM) to resolve the nature of HER2 nanocluster on the cell surface. Single molecule localizations showed that HER2 mainly exists as small nanoclusters in cells treated with NC antibody. In zani treated cells HER2 molecules coalesce to form large spatially-separated clusters. This is confirmed by clustering and mean area analysis of images from multiple regions of interest. Though cells stimulated with tras, pert and tras+pert displayed varying degrees of HER2 organization, large nanoclusters observed with zani were absent in cells treated with these agents.

**Conclusion**

- The biparatopic design of zanidatamab results in a unique networked complex geometry upon binding to HER2.
- Structural modeling of zanidatamab with HER2 supports its ability to form complexes with distinct higher order geometry on target cell surfaces, potentially promoting novel function such as CDC (Complement Dependent Cytotoxicity) that is not observed with either tras, pert or their combination (See Abstract number 1005).
- High resolution imaging of HER2 organization on cells supports the structural modeling studies, confirming the presence of unique large cluster formations not seen with tras, pert or their combination.
- The observed differences in the geometry, size and persistence of these coalesced cell surface receptor clusters likely contribute to the differentiated activity of zanidatamab.

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