Short Peptide Nucleic Acid-IGF1 Tetrapeptides Enable Specific MicroRNA Blockade in Triple Negative Breast Cancer Cells without Passenger Strand Side Effects

252nd American Chemical Society National Meeting



Bound Therapeutics LLC

Breast cancer subtypes



http://www.pathophys.org/breast-cancer/

OncomiRs in cancer



Iorio, M. V. and C. M. Croce (2012), EMBO Mol Med 4(3): 143-159

miRNA Biogenesis and Mechanisms of Action



Iorio, M. V. and C. M. Croce (2012), EMBO Mol Med 4(3): 143-159

AKT activation is an interplay between miR-21 and miR-17.



miRNA inhibition by modified oligonucleotides



miR-17-5p knockdown by DNA-LNA chimera unexpectedly decreased PDCD4 and PTEN protein in MDA-MB-231 cells.



Jin, Y. Y., et al. (2015), PLoS One 10(12): e0142574

pre-miRNA structure of miR-17 revealed sequence similarity between DNA-LNA chimera and miR-17-3p passenger strand.



5' <mark>A-CUGCA</mark>G<mark>UG-AAGGCAC-UUG</mark>UAG 3' miR-17-3p 5' <mark>ACCTGCA</mark>C<mark>TG</mark>TAAG-CACTTTG 3' Anti-miR-17-5p LNA

Jin, Y. Y., et al. (2015), PLoS One 10(12): e0142574

Competition between anti-miR-17-5p and miR-17-5p for inhibition of *PDCD4* mRNA



Jin, Y. Y., et al. (2015), PLoS One 10(12): e0142574

Luciferase assay system to test anti-miR-17 – mRNA interaction





Anti-miR-17-5p DNA-LNA lowered the expression of luciferase vectors containing several predicted *PDCD4* and *PTEN*'s 3'UTR target sites for miR-17-3p.



Jin, Y. Y., et al. (2015), PLoS One 10(12): e0142574

miRNA blocker design strategy

- Eliminate extra side-effects of conventional microRNA blockers
- TNBC cell-specific delivery method
- No complicated formulation, soluble in saline, intravenous route
- Next generation RNA backbones (FANA & NC-BNA vs. PNA) will elevate efficacy and potency



Jin *et al*. (2015) *PLoS One* **10**:10.1371/journal.pone.0142574

Nucleotide Analog - Peptide Nucleic Acids Increasing stability, binding affinity and specificity

- High binding affinity to complementary DNA/RNA.
- Differentiation of single-base mismatch by high destabilizing effect.
- High chemical stability to temperature and pH.
- High biological stability to nuclease and protease.
- Good uptake via basic peptides or receptor-specific ligands
- Mice given up to 100 mg/kg dose of PNA-peptide conjugate daily did not show any irreversible toxicity (Chaubey et al., 2008).



Chaubey, B., et al. (2008), Oligonucleotides **18**(1): 9-20 Good & Nielsen (1997) *Antisense Nucl Acid Drug Dev.* **7**: 431-437

Delivery - IGF1 retro-inverso analog



In vivo specificity of 12-mer PNA-IGF1 tetrapeptides

External Imaging of CCND1 Cancer Gene Activity in Experimental Human Breast Cancer Xenografts with ^{99m}Tc-Peptide-Peptide Nucleic Acid-Peptide Chimeras

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24 hr

In vivo specificity of 12-mer PNA-IGF1 tetrapeptides



Tian, X., et al. (2007), J Nucl Med 48(10): 1699-1707

MDA-MB-231 cell uptake of Cal560-Anti-miR PNA-IGF1 tetrapeptide



Cells were incubated in 200 nM of Cal560-Anti-miR PNA-IGF1 tetrapeptide and negative controls for 4 hours at 37°C in complete medium. Ex: 543 Em: 560

μ M anti-miR PNA-IGF1 tetrapeptide elevated the expression of PDCD4 and PTEN



Fig. 1. PNA-AEEA-cyclo-D(Cys-Ser-Lys-Cys) blocker of miR-17-5p.



Blocking miR-21 with anti-miR-21 PNA-IGF1 tetrapeptide slowed down MDA-MB-231 cell migration.



Blocking miR-21 with PNA-IGF1 tetrapeptide slowed down MDA-MB-157 cell migration.





Blocking miR-21/17 with PNA-IGF1 tetrapeptide induced apoptosis in MSL type MDA-MB-231 and MDA-MB-157 cells.



Summary

- The functional changes as a result of 1 μM PNA-IGF1 peptide treatment are modest, indicating low efficacy.
- TNBC cells that rely on PI3K/AKT/mTOR pathway are likely to respond to miR-21/17 blockage.
- Future antagomiRs can be optimized by:
 - Alternative oligonucleotide analog that triggers RNase H (NC-BNA, FANA)



- Increasing the length of antagomiRs without mimicking passenger strand
- Better delivery target

Yamamoto, T., et al. (2012), Mol Ther Nucleic Acids **1**: e22 Kalota, A., et al. (2006), Nucleic Acids Res **34**(2): 451-461

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