

Introduction

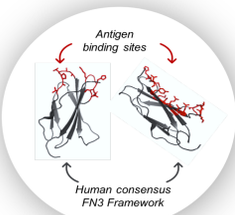
Aro Biotherapeutics is a preclinical stage biotechnology company focused on discovery and development of Centyrins, a new class of small, structurally simple, highly stable and soluble proteins engineered to specifically bind antigens with high affinity.

We demonstrate Centyrins have a longer residency time in the early endosome relative to antibodies which are rapidly shunted to lysosomes after binding to the same receptor. Using Centyrins targeted to cell surface receptors on tumor cells, we also demonstrate efficient internalization and trafficking of Centyrins to the cytosol via protein complementation as demonstrated using GFP complementation assays.

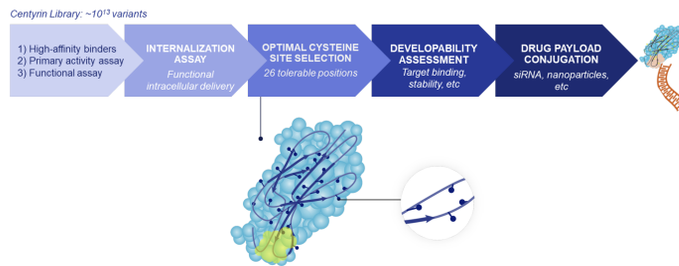
Centyrins provide a means to specifically deliver oligonucleotides to cell types beyond hepatocytes which enables access to intracellular targets that have been considered "undruggable". Our Centyrin-KRAS siRNA conjugates are designed to inhibit a variety of solid tumors driven by KRAS mutations and are released in early endosomes over an extended time period. These data highlight the broad utility of this platform.

Centyrins for Delivering siRNA

Small interchangeable protein scaffolds, optimized for multi-antigen targeting and delivery, of complex drug payloads, including RNA drugs



Small and Simple	Excellent tissue and tumor penetration; cleared through kidney No glycosylation sites Site specific drug conjugation
Unique Biophysical Properties	Soluble, heat and pH stable E coli production with high expression levels
Flexible	Rapid screening of Centyrin libraries using <i>in vitro</i> CIS-display and intracellular delivery Centyrins selectively bind proteins with high affinity and specificity Easily formatted into bi- and multi- specific molecules



Centyrins Reside in Endosomes Longer than Antibodies

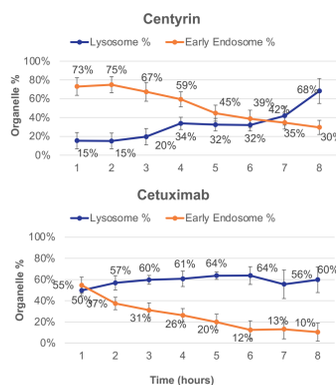
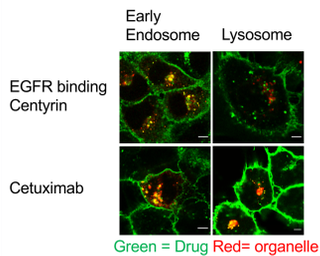


Figure 2 : CAL 27-SCCHN cells were transiently transfected with LAMP1 or Rab5a rp and were treated with an EGFR targeting Centyrin or Cetuximab labeled with Alexa-Fluor 488.

Figure 3 : Single cell imaging was used to track the trafficking of each construct. An average of 10 cells were used for each point.

Centyrins Traffic into the Cytoplasm

"Split GFP" experiment

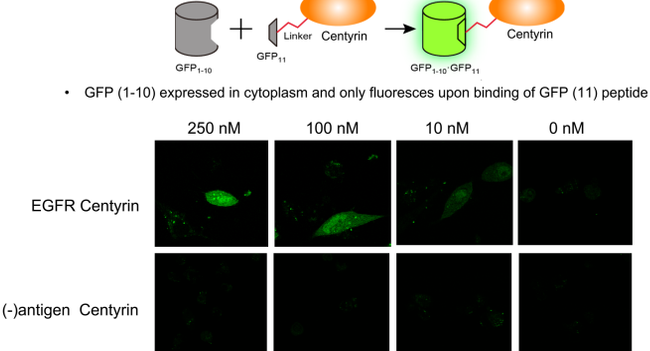


Figure 4 : An EGFR binding Centyrin and a control (-) antigen negative Centyrin were expressed with GFP11 amino acid sequence at the C-terminus. HCC872 cell were treated for 24 h with each Centyrin and then imaged.

Targeting Various Mutants of KRAS

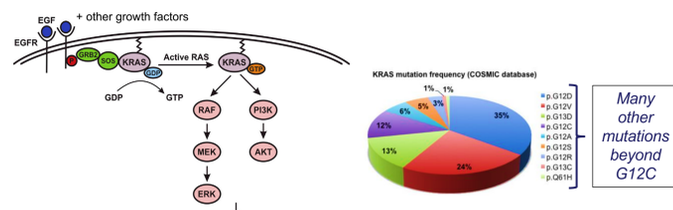
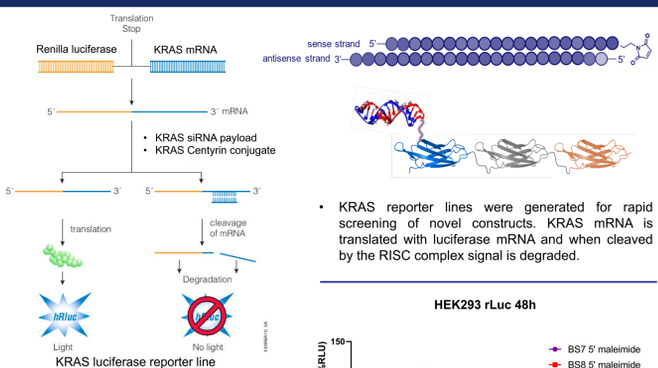


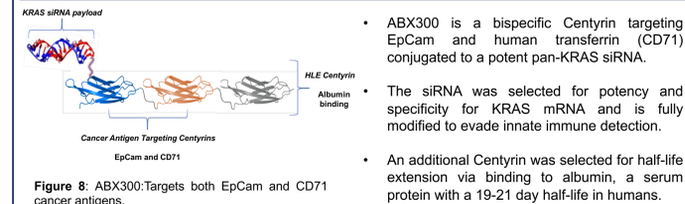
Figure 5 : KRAS is a critical regulator of the MAPK/ERK pathway responsible for tumor oncogenesis in solid many tumors. **Figure 6** : While therapies are emerging for G12C, ~12 % of mutant forms, other KRAS mutant cancers are still undruggable.

siRNA Sequence/Chemistry SAR



- Over 3500 siRNA sequences were computationally evaluated for off target effects and potency against pan-KRAS mRNA.
- 87 siRNAs were tested as fully modified and protected sequences via transfection into HEK293 luciferase based reporter cell line.
- The most potent and selective sequences were further modified with maleimide linker chemistry to enable conjugation to the targeting Centyrin, Figure 7.

ABX300: Dual Antigen Targeting



- ABX300 is a bispecific Centyrin targeting EpCam and human transferrin (CD71) conjugated to a potent pan-KRAS siRNA.
- The siRNA was selected for potency and specificity for KRAS mRNA and is fully modified to evade innate immune detection.
- An additional Centyrin was selected for half-life extension via binding to albumin, a serum protein with a 19-21 day half-life in humans.

Advantages of dual targeting

- Avidity increases internalization rate and accumulation of therapeutic payload.
- Dual antigen specificity further differentiates targeting to tumor specific tissue.
- Ease of Centyrin production allows for rapid screening of target combinations for the most effective combination of targets.

ABX300 Inhibits Cancer Cell Proliferation

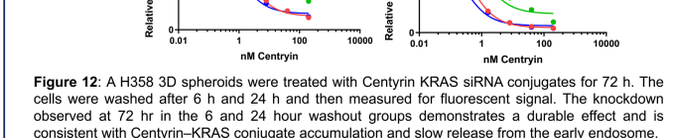
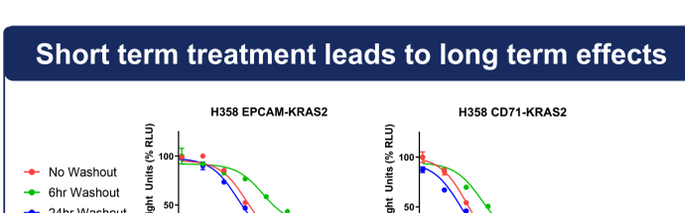
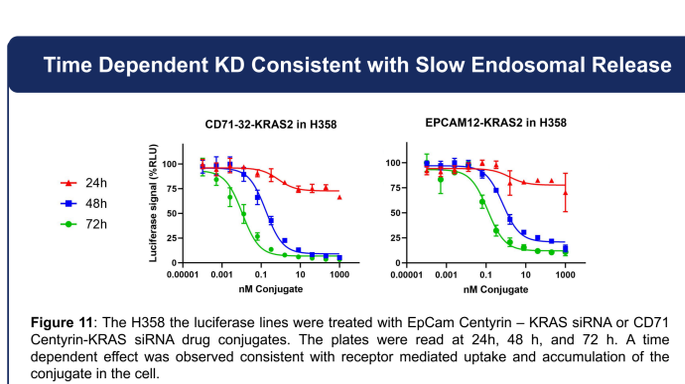


Figure 11 : The H358 the luciferase lines were treated with EpCam Centyrin – KRAS siRNA or CD71 Centyrin-KRAS siRNA drug conjugates. The plates were read at 24h, 48 h, and 72 h. A time dependent effect was observed consistent with receptor mediated uptake and accumulation of the conjugate in the cell. **Figure 12** : A H358 3D spheroids were treated with Centyrin KRAS siRNA conjugates for 72 h. The cells were washed after 6 h and 24 h and then measured for fluorescent signal. The knockdown observed at 72 hr in the 6 and 24 hour washout groups demonstrates a durable effect and is consistent with Centyrin–KRAS conjugate accumulation and slow release from the early endosome.

ABX300 induces KD on G12C and other KRAS mutants cell lines

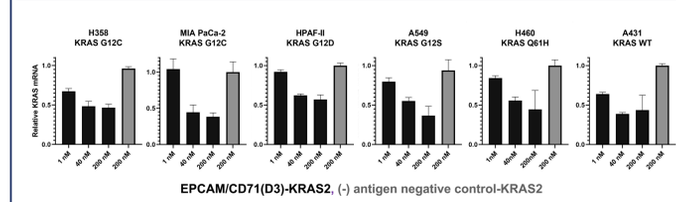


Figure 13 : H358-NSCLC (G12C), MIA PaCa-2-pancreatic (G12C), HPAF II-pancreatic (G12D), A549-NSCLC (G12S), H460-NSCLC (Q61H) and A431-skin (KRAS WT) cancer lines were treated with Centyrin-KRAS conjugates for 72 h. The cells from each experiment were measured for residual KRAS mRNA using qPCR.

ABX300 Decreases KRAS Protein Levels

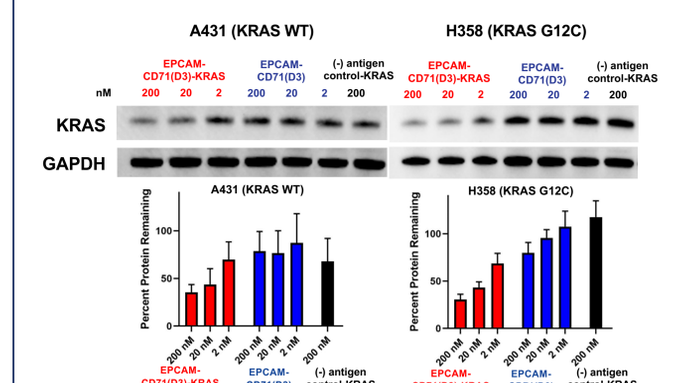


Figure 14 : A431 and H358 cells were treated with bispecific Centyrins conjugated to KRAS2 at 2, 20 and 200 nM concentrations. Western blots were used to detect KRAS protein at 72 hours. A good correlation between siRNA mRNA silencing and protein was observed.

Conclusions

- ABX300 demonstrated receptor specific delivery of KRAS siRNA.
- Centyrin:siRNA conjugates demonstrated high potency against tumor cell lines.
- Centyrins possess differentiated trafficking vs. antibodies facilitating siRNA delivery.
- Rapid generation of Centyrin combinations allowed for the selection of potent EpCam and CD71 Centyrin combinations for delivery of any siRNA payload into tumor cells.
- Potential broad utility of Centyrins to deliver siRNA or other payloads into many internalizing receptor positive cells.

References

1. Hobbs, A.; Der, C.; Rossman, K. "RAS isoforms and mutations in cancer at a glance" *J. Cell. Sci.*, 2016, 1287.