I. Choosing a target sequence

A. Client chooses a target gene and submits to the Core a CRISPR Request Form that includes: Client’s name and email, PI name, account to be charged, gene name, and sequences of two ‘potential target regions’ of the gene.

Criteria for ‘potential target regions’:
- Preferred minimum size of region: 200bp
- Often the two ‘potential target regions’ would be two different exons. One large exon can be submitted instead of two separate regions. To create a null allele it is best to target an exon that is as 5’ in the gene as possible, and certainly that is 5’ of a sequence encoding a conserved portion of the protein (for example upstream of a DNA binding domain).
- If only small exons are available, please submit the exon along with 20bp of upstream and downstream intron sequence
- We normally avoid exon 1 to avoid possible alternative transcription start sites, but if these can be ruled out by the client it is ok to target exon 1
- Beware of alternatively spliced exons

B. Core identifies 2-3 best candidate CRISPR target sequences and sends this information to Client.

C. Client then chooses one of the candidate target sequences and verifies that their model organism to be used has the exact CRISPR-binding sequence without polymorphisms.

One approach to doing this in Zebrafish is:
- Fin clip and prepare genomic DNA from a series of adults. Amplify from one gDNA sample 300 – 600 bp of genomic sequence that includes the target site. Sequence the amplicon. If the sequence matches the candidate target, you will analyze more adults. If the sequence does not match the candidate target, try a different WT strain.
- Genotype additional adult fish (4 - 8 from each sex from the same strain). This can be accomplished by either of two strategies: 1) sequence additional adults; 2) use High Resolution Melt Analysis to determine if the additional adults have any signs of polymorphisms in the region. We highly recommend preparing HRMA primers, as they will likely be used to detect mutations. It is advisable to design the sequencing primers above so that one of the two sequencing primers can also be used for HRMA assays (HRMA primers should generate a 75 – 120bp amplicon that includes the entire target sequence). Once you find adults that carry the target sequence, the target sequence is considered verified.
- Once the target site is verified, contact the Core to request CRISPR gene construction against the target. The Core will not begin construction until the target is verified.
II. CRISPR gene construction

The Core will generate two plasmids, encoding the Custom sgRNA for your target site and encoding the Cas9 endonuclease. Constructions are sequence-verified and should be completed in about one week.

III. Optimizing HRMA conditions to detect polymorphisms

The Client needs to prepare primers for HRMA and be able to detect newly induced polymorphisms at the target site. Please contact the Core (mutrus@genetics.utah.edu) for any help with the design or implementation of HRMA detection of polymorphisms.

IV. Feedback to the Core

We need to monitor how well our methods are working. This is a new technology, and it is still being optimized! We rely on your help! Upon testing the induction of mutations, please contact the Core and report to us:

- Gene name and CRISPR name, # injected embryos with evidence of new mutations / # injected embryos assayed. Please ask for advice and give us feedback. We are here to be helpful.