



# VCU

## Office of Research and Innovation **spotlights...**

September 2015

### VCU Transgenic/Knockout Mouse Core

The mission of the VCU Transgenic/Knockout (TG/KO) Mouse Core is to provide an efficient and economical means for producing genetically modified mice, and to provide support services to facilitate the use of these models by VCU researchers.

#### Genetically modified mice include:

- Transgenic mice, in which customized genes are introduced into the mouse genome (either by random integration or by targeted insertion into a pre-determined locus)
- Knockout mice, in which an endogenous mouse gene has been inactivated
- Knock-in mice, in which an endogenous mouse gene has been modified (e.g., "floxed", introduction of point mutation, etc)

#### Additional TG/KO Core services include:

- Breeding colony management
- Genotyping
- Mouse line rederivation
- Sperm and embryo cryopreservation
- MMRB IVIS imager management
- Consulting on project design and all things mouse-related

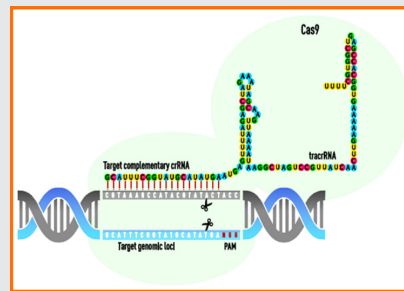
#### INTRODUCING CRISPR/CAS9 FOR GENOME EDITING!

Until recently, most knockout and knock-in mice were made by the same laborious methods originally developed in the mid-1980s. In late 2013, all that changed with the advent of CRISPR/Cas9, a system for creating RNA-guided DNA double strand breaks at precise genomic locations (repurposed from a bacterial adaptive immune defense against phage infection). The use of CRISPR/Cas9 technology eliminates the need for targeting in ES cells and generating chimeras – and makes KO mouse production as easy as generating transgenic mice!

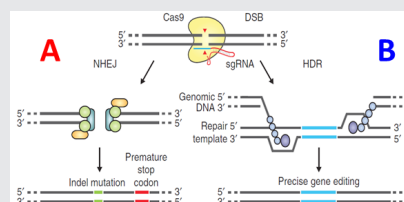
The TG/KO is now offering CRISPR/Cas9 services, including:

- Design and synthesis of sgRNA and design of DNA repair templates (ssODNs)
- Microinjection of fertilized mouse eggs and screening of resulting pups for presence of the desired allele
- Screening for off-target mutation events

Contact Jolene Windle to discuss your CRISPR/Cas9 project!



Courtesy of Life Technologies™



**Figure 2** | DSB repair promotes gene editing. DSBs induced by Cas9 (yellow) can be repaired in one of two ways. In the error-prone NHEJ pathway, the ends of a DSB are processed by endogenous DNA repair machinery and rejoined, which can result in random indel mutations at the site of junction. Indel mutations occurring within the coding region of a gene can result in frameshifts and the creation of a premature stop codon, resulting in gene knockout. Alternatively, a repair template in the form of a plasmid or ssODN can be supplied to leverage the HDR pathway, which allows high fidelity and precise editing. Single-stranded nicks to the DNA can also induce HDR.

Nature Protocols 8:2281-2308, 2013

#### Minimal requirements:

- a custom "sgRNA" consisting of both guide RNA (homologous to desired cleavage site) and scaffold RNA sequences
- Cas9 enzyme
- Optional: a single-strand or double-strand DNA repair template

These are co-injected into fertilized mouse eggs.

**A** Following creation of a double-strand break, the cell can rejoin the two ends by **Non-Homologous End Joining**, often creating small insertions or deletions. If this is in the coding region, it may create a frameshift that results in a **KO allele**.

– OR –

**B** If a DNA template homologous to the cleavage site but containing a desired modification (e.g., point mutation, loxP site, etc) is included, **Homology-Directed Repair** may create a **modified allele** containing the desired modification.

**BOTH PROCESSES ARE HIGHLY EFFICIENT!**



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