

Transgenes (TG)

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| Basic Vector TG | Can target anywhere |
| TG with some homology | Target to least homologous region |
| Promoter-- Vector TG | Target at promoter-vector junction |
| Mouse cDNA | Target at exon-exon junction (Be aware of intron size) |
| Human TG | Target to least homologous region |

SNPs (MUT)

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| Basic SNP | Center SNPs in reporter oligos. |
| SNP based off AA change given | Locate AA change. Center SNPs in reporter oligos. |
| rs# | Locate rs#. Ensure reporter 1 is BL6 sequence. Center SNPs in reporter oligos. |

Both reporters MUST be on the same strand or the assay will not work.

Humanized Models (KO)

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| Standard humanized KO model | Identify endogenous human junction and include 2-3 mismatches per oligo (WT and Mutant) if possible. |
| Humanized model with FRT or LoxP | Target flanking the FRT/LoxP which allows you to avoid homology |

Knockouts (KO)

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| Standard KO Model | Target KO assay at 5' or 3' junction |
| Small CRISPR deletion | Balance KO probe across new junction |
| Large CRISPR deletion | Flank new junction |

Designs for corresponding WT will be noted in the wildtype section.

KO landmarks

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| Start/Stop Region | Design WT at start or stop (depending on model). KO probe will be a generic. |
| Coding Region | Design anywhere in coding region. KO probe will be a generic |
| Restriction Site | Design according to restriction sites. KO probe will be a generic. |
| Coordinates | Design according to coordinates. KO probe will be a generic. |

When in doubt - SEQUENCE.

Wildtypes (WT)

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| WT for small insertion (<400bp) | Balance probe (40/30) or do allele specific primer. |
| WT for large insertion (>400bp) | Can flank insertion site or have a probe across insertion site but do NOT need to balance. |
| WT where some bases were deleted (either via CRISPR or upon insertion of vector) | Target probe within deleted bases. Primers can be on either side. |

Floxed (FL or MD)

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| Floxed allele with vector | Flank LoxP, use tearpin, or break the hairpin. |
| Floxed allele with minimal vector | Use tearpin or break the hairpin |
| Floxed allele when unsure which LoxP | Flank LoxP or do tearpin |
| Marker deleted floxed allele | Flank FRT/LoxP |

Identify which LoxP you have. If you are breaking the hairpin you MUST know which LoxP to ensure you break in the correct direction.

Excised (EX)

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| EX with vector | Flank remaining LoxP or tearpin. |
| EX with minimal vector | Tearpin or can flank LoxP if large enough deletion created by Cre recombination. |

CRISPR

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|------|---|
| NHEJ | Flank new junction created as long as it is >400bp |
| HDR | Must target 1 oligo outside of ssODN to be site specific. |

Remember that CRISPR rules can apply to a variety of models including SNPs, Conditional lines, and even traditional KOs. Please be aware if you are designing for CRISPR and utilize our best practices.



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