Harnessing CRISPR to boost NGS sensitivity with CRISPRclean™

The revolutionary power of the CRISPR system

The CRISPR gene editing system has revolutionized biological and medical research. CRISPR consists of the Cas9 enzyme that targets DNA for cleavage in a site-specific manner when complexed with CRISPR guide RNA (gRNA) (Figure 1). Originally evolved to degrade viral genomes, the CRISPR system was later modified by scientists to become an important molecular tool. The adaptability of the CRISPR system enabled researchers to precisely target locations in the genome in a broad range of applications, such as gene editing, gene therapy, functional library screening, and more.

CRISPRclean technology harnesses the specificity of CRISPR to refine NGS libraries

Like CRISPR, next-generation sequencing (NGS) technology has also revolutionized the research world. The ability to sequence genomes, transcriptomes, and epigenomes without a priori knowledge has yielded an unprecedented amount of discoveries about genes and genetic variation. As NGS technology continues to advance with innovations to increase sensitivity and throughput while lowering costs, analysis of previously elusive samples are possible with unprecedented depth, such as single cell transcriptomes, or complex microbial communities.

Although NGS provides millions of reads from a given sample, much of the sequencing information can be biologically uninformative. This is particularly the case for RNA sequencing when ribosomal RNA (rRNA) comprises up to 90% of total RNA in both eukaryotes and prokaryotes. Thus, refining the sample to focus on more informative sequences can increase sensitivity by lowering background from abundantly expressed genes and may also save significantly on sequencing costs.

This technology is integrated into a stranded total RNA sequencing library prep protocol by adding Cas9/RNA complexes after the adapter ligation step. The Cas9/RNA complexes consist of a pool of guide RNAs that target and deplete unwanted sequences in each application, but the CRISPR mechanism does not rely upon specific sequencing platforms.

Legacy targeted resequencing approaches trade off the risk of introducing bias with the benefit of improved sensitivity of desired genes or transcripts. CRISPRclean differs from these methods by performing negative selection of specific abundant, uninformative sequences, thus maintaining the ability to analyze informative sequences without a priori knowledge of which transcripts or variants will be relevant in a given study. Instead of fishing for a needle in a haystack, CRISPRclean removes the haystack, enabling analysis of the remaining “needles.” This results in increased sequencing coverage across genes or transcripts of interest that remain in a library.

CRISPRclean technology is both specific and highly programmable towards unwanted sequences in any given application. Beyond ribosomal RNA depletion,
CRISPRclean can be designed to deplete unwanted sequences from virtually any application. For example, a range of highly expressed genes that are relevant to specific tissues can be depleted during single-cell analysis. For DNA applications, CRISPRclean technology can target repetitive elements or other undesired wild-type loci.

Other key features of CRISPRclean technology add further flexibility towards a broad range of applications. By using negative selection, in which the targeted “haystack” removal approach only requires information about targeted sequences, CRISPRclean technology retains the ability to discover new variants of low abundance with its agnostic approach. Furthermore, the post-library depletion workflow has enabled CRISPRclean to access low input samples while maintaining performance, thus demonstrating its utility for samples from which nucleic acids are difficult to obtain.

The CRISPRclean workflow

CRISPRclean ribodepletion (Figure 3) is integrated into a stranded RNA library prep protocol. After targeted sequences are cut, they cannot be substrates for PCR amplification and subsequent sequencing. The refined libraries (non-targeted sequences) is size-selected using magnetic beads, and then amplified a small number of rounds. The output is a refined, sequence-ready NGS library.

Several approaches to DNA analysis involve enrichment for selected genes or loci by positive target selection, either through primer-directed amplification or oligo-based hybrid capture. By performing negative selection of specific selected sequences, CRISPRclean technology maintains an unbiased approach to analysis of informative sequences that may include undiscovered variants.

Figure 2: CRISPRclean mediated depletion fits into a stranded RNA library prep protocol integrating seamlessly into NGS workflows. After degradation of unwanted sequences, informative, lower abundance sequences are refined for sequencing.
Benefits of CRISPRclean technology

- Highly programmable
- Achieves efficient depletion
- Increased sensitivity
- Simple automatable workflow

Summary

CRISPRclean is a novel method for depletion of abundant or uninformative sequences in NGS libraries. It can increase the ability to detect rare transcripts in RNA applications, or analyze low-abundance variants in DNA applications, by increasing depth of coverage of informative sequences. The reduction of high background associated with abundant sequences further adds confidence to sequencing data for detection of rare sequences. Furthermore, focusing sequencing platforms on the most relevant information in genomic and transcriptomes can significantly reduce costs.

Examples of applications which CRISPRclean technology can be applied to are ribosomal removal, metatranscriptomics, infectious disease pathogen detection, host immune response profiling, single cell analysis, and more.

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Catalog

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Figure 3: CRISPRclean Stranded Total RNA Prep with rRNA Depletion (Human, Mouse, Rat) is a 1-day workflow comprised of 7 steps and approximately 3 hours hands-on time.