

LASERGENE VERSION COMPARISON

Lasergene Molecular Biology	15	16	17
Pairwise and Multiple Sequence Alignment			
Variant analysis for viral, mitochondrial, and chloroplast genomes New in Lasergene 17-4			•
MAFFT7 algorithm for faster alignments of viral genomes and 16S data sets			•
High-capacity, customizable distance tables that support large data sets			•
Improved alignment report that can be configured and exported			•
RAxML, for Maximum Likelihood phylogenetic inference calculation			•
Support for multiple phylogenetic trees in a single alignment			•
Perform BLAST and Entrez searching from MegAlign Pro alignments		•	•
Updated views and enhanced user controls in MegAlign Pro		•	•
Export editable images to PowerPoint	•	•	•
Profile alignment functionality to merge alignments or additional sequences	•	•	•
Sanger Sequence Assembly and Analysis			
Full integration of SeqMan NGen into SeqMan Ultra for streamlined assembly			•
Easily export an image of your assembly in PDF, BMP, or PPT format			•
Improved vector trimming with extensive, built-in vector database			•
Easy multiple-sample variant analysis and visualization			•
New 64-bit SeqMan Ultra application with better performance and increased capacity			•
Assess read alignment, coverage, and SNPs	•	•	•
Assemble reads de novo or against one or more reference sequences	•	•	•
Cloning, Primer Design, and Gene Detection			
Batch cloning to automatically clone one or more fragments simultaneously			•
Enhanced support for VectorNTI, Geneious, and Clone Manager Suite file types		•	•
PCR site-directed mutagenesis with ability to predict impact of mutation on protein structure		•	•
Agarose gel simulation functionality	•	•	•
Plasmid auto-annotation (single or batch) using curated database	•	•	•
Automated virtual cloning: site-directed, TA, TOPO, Gateway, InFusion, GeneArt, and Gibson Assembly	•	•	•
PCR primer design	•	•	•

Add our Genomics and Protein Applications to Complete Your Lasergene Package! See reverse side for details.

Lasergene Genomics	15	16	17
Next-Gen Sequence Assembly and Alignment			
Expanded viral genome assembly to support data from ARTIC protocols			•
Support for <i>de novo</i> genome assembly with PacBio HiFi reads			•
Full integration of SeqMan NGen into SeqMan Ultra for streamlined assembly			•
Unique, mutli-sample visualization to view and compare assembled sequences			•
Support for combining multiple projects of the same type into a single project			•
64-bit SeqMan Ultra application for project analysis with better performance and increased capacity			•
Auto-analysis of hardware and data to determine if assembly should be run locally or on the Cloud			•
Ability to polish Canu or Spades assemblies of PacBio and Oxford Nanopore long read data			•
miRNA quantitation workflow to quantitate and analyze miRNA gene expression levels			•
Integrated DNASTAR Cloud Assemblies for projects that exceed the capacity of your hardware		•	•
Quick access to NGS and Sanger project setup in the new DNASTAR Navigator		•	•
Enhanced RNA-Seq statistics with DESeq and edgeR from Bioconductor	•	•	•
RNA-Seq analysis for model and non-model organisms	•	•	•
Gene panel workflows with control validation	•	•	•
De novo genome and transcriptome assembly	•	•	•
Variant Detection and Analysis			
Mastermind integration to cross-reference variant data in genomic literature			•
Updated Variant Annotation Database includes dbNSFP version 4.1			•
Live variant report updates variants in real-time, as filters are applied			•
Ability to compare and analyze multiple VCF files from other NGS software pipelines			•
Model detected variants on protein structure with updated protein design workflow*		•	•
Combine variant and PDB annotations to predict potential effects on protein structure*	•	•	•
Combine SNP and CNV analysis with any workflow	•	•	•
Direct comparison to dbSNP and GERP and dbNSFP databases	•	•	•
Lasergene Protein	15	16	17
Macromolecular Structure, Motion, and Function			
Ability to import UniProt files, with full support for displaying annotations			•
Support for exporting data used to generate Analysis View tracks			•
Protein design tools to improve fold stability and developability*		•	•
Integrated BLAST and Entrez searching in Protean 3D		•	•
Mutation modeling and neighbor search to analyze SNP impact on protein structure	•	•	•
Access to NovaDock protein-protein docking, analysis and visualization*	•	•	•
Access to NovaFold and NovaFold Antibody structure prediction*	•	•	•
Protein function and ligand binding site prediction*	•	•	•
B-cell epitope prediction	•	•	•
* Additional subscription may be required to access protein modeling and design programs			

