■ module 経由で使えるもの

事前に設定ファイルを読み込む必要があります

source /apl/bio/etc/bio.sh

Analysis Type	Application name	Description	installed ver.	Official URL		
Homology search	blast+	Sequence Similarity Search	2.16.0	https://blast.ncbi.nlm.nih.gov/Blast.cgi		
	blat	BLAT on DNA is designed to quickly find sequences of 95% and greater similarity of length 25 bases or more	https://genome.ucsc.edu/goldenpath/help/blatSpec.html			
	Diamond	DIAMOND is a sequence aligner for protein and translated DNA searches, designed for high performance analysis of big sequence data.	2.1.11	https://github.com/bbuchfink/diamond		
	fasta	Sequence Similarity Search	36.3.8g			
	HH-suite	The HH-suite is an open-source software package for sensitive protein sequence searching based on the pairwise alignment of hidden Markov models (HMMs).	3.3.0_SSE2	https://github.com/soedinglab/hh-suite		
	MMseq2	ultra fast and sensitive sequence search and clustering suite	17-b804f	https://github.com/soedinglab/MMseqs2		
	vsearch	VSEARCH is an open source and free of charge multithreaded 64-bit tool for processing metagenomic nucleotide sequence data. An alternative to the USEARCH	2.29.0	https://github.com/torognes/vsearch		
	bamtools	Bamtools is a toolkit for analyzing and managing BAM files	2.5.2	https://github.com/pezmaster31/bamtools		
	bedops	BEDOPS: the fast, highly scalable and easily-parallelizable genome analysis toolkit	2.4.41	https://bedops.readthedocs.io/en/latest/index.html		
	BEDtools	Bedtools utilities are a tools for a wide- range of genomics analysis tasks	2.31.1	https://bedtools.readthedocs.io/en/latest/		
	Bowtie	Bowtie is an ultrafast, memory-efficient short read aligner	1.2.3	https://bowtie-bio.sourceforge.net/index.shtml		
	Bowtie2	Bowtie 2 is an ultrafast and memory- efficient tool for aligning sequencing reads to long reference sequences.	2.5.3	https://bowtie-bio.sourceforge.net/bowtie2/index.shtml		
	bwa bwa-mem	Burrows-Wheeler Aligner (BWA) is an efficient program that aligns relatively short nucleotide sequences against a long reference sequence	0.17.7 2.2.1	https://github.com/lh3/bwa		
	Cufflinks	Cufflinks assembles transcripts, estimates their abundances, and tests for differential expression and regulation in RNA-Seq samples		https://github.com/cole-trapnell-lab/cufflinks		
	fastQC	A quality control tool for high throughput sequence data.		https://www.bioinformatics.babraham.ac.uk/projects/fastqo		
	fastp	A quality control tool for high throughput sequence data.	0.23.4	https://github.com/OpenGene/fastp		
	GATK	The GATK is the industry standard for identifying SNPs and indels in germline DNA and RNAseq data.				
	hisat2	HISAT2 is a fast and sensitive alignment program for mapping next-generation sequencing reads (both DNA and RNA).	2.2.1	https://daehwankimlab.github.io/hisat2/		
	kallisto	kallisto is a program for quantifying abundances of transcripts from RNA-Seq data, or more generally of target sequences using high-throughput sequencing reads.	0.46.2	https://github.com/pachterlab/kallisto		
	MACS2	Model-based Analysis of ChIP-Seq on short reads sequencers such as Genome Analyzer (Illumina / Solexa)	2.1.2	https://github.com/macs3-project/MACS		
	RSEM	RNA-Seq by Expectation-Maximization	1.3.3	https://github.com/deweylab/RSEM		
	Salmon	Salmon is a tool for quantifying the expression of transcripts using RNA-seq data	1.10.0	https://combine-lab.github.io/salmon/		

Application name	Description	installed ver.	Official URL		
samtools	SAM Tools provide various utilities for manipulating alignments in the SAM format	1.19.2	https://www.htslib.org/		
seqkit	SeqKit - a cross-platform and ultrafast toolkit for FASTA/Q file manipulation	2.10.0	https://github.com/shenwei356/seqkit		
soap	Short Oligonucleotide Analysis Package	2.21	https://github.com/ShujiaHuang/SOAPaligner		
SRAtoolkit	The SRA Toolkit and SDK from NCBI is a collection of tools and libraries for using data in the INSDC Sequence Read Archives.	3.0.0	https://github.com/ncbi/sra-tools		
STAR	Spliced Transcripts Alignment to a Reference	2.7.11b	https://github.com/alexdobin/STAR		
Stringtie	StringTie is a fast and highly efficient assembler of RNA-Seq alignments into potential transcripts.	3.0.0	https://ccb.jhu.edu/software/stringtie/		
Tophat	TopHat is a fast splice junction mapper for RNA-Seq reads	2.1.1	https://ccb.jhu.edu/software/tophat/index.shtml		
CAT_pack	A pipelines for the taxonomic classification of long DNA sequences and metagenome assembled genomes (MAGs / bins)	6.0.1	https://github.com/MGXlab/CAT_pack		
ABySS	Assembly By Short Sequences - a de novo, parallel, paired-end sequence assembler	2.3.4	https://github.com/bcgsc/abyss		
Allpaths-LG	The new short read genome assembler.	52488	https://www.rcac.purdue.edu/software/allpathslg		
canu	Canu is a fork of the Celera Assembler designed for high-noise single-molecule sequencing (such as the PacBio RSII or Oxford Nanopore MinION).	2.2	https://github.com/marbl/canu		
hifiasm	Hifiasm is a fast haplotype-resolved de novo assembler for PacBio HiFi reads.	0.18.5	https://github.com/chhylp123/hifiasm		
MaSuRCA	MaSuRCA (Maryland Super-Read Celera Assembler) genome assembly software	4.0.7	https://github.com/alekseyzimin/masurca		
NECAT	NECAT is an error correction and de-novo assembly tool for Nanopore long noisy reads.	0.0.1	https://github.com/xiaochuanle/NECAT		
SPAdes	SPAdes St. Petersburg genome assembler: SPAdes is an assembly toolkit containing various assembly pipelines.	4.1.0	https://github.com/ablab/spades		
Trinityrnaseq	Novel method for the efficient and robust de novo reconstruction of transcriptomes from RNA-seq data	2.15.1	https://github.com/trinityrnaseq/trinityrnaseq		
velvet	Sequence assembler for very short reads	1.2.10	https://github.com/dzerbino/velvet		
soap denovo	Short Oligonucleotide Analysis Package genome assembler	2.04	https://github.com/ShujiaHuang/SOAPaligner		
wgs	same as Celera Assembler: Whole genome assembler	8.2	https://github.com/alekseyzimin/wgs		
lastz	A tool for (1) aligning two DNA sequences, and (2) inferring appropriate scoring parameters automatically	1.04	https://github.com/lastz/lastz		
MUMmer	MUMmer is a system for rapidly aligning entire genomes, whether in complete or draft form.	4.0.0	https://github.com/mummer4/mummer		
clustal Omega	Fast, accurate, scalable multiple sequence alignment for proteins	1.2.4	http://www.clustal.org/omega/		
clustalw	Multiple Sequence Alignment	1.83	http://www.clustal.org/clustal2/		
clustalw2	Multiple Sequence Alignment	2.1	http://www.clustal.org/clustal2/		
FAMSA	Progressive algorithm for large-scale multiple sequence alignments.	1.6.2	https://github.com/refresh-bio/FAMSA		
Gblocks	Gblocks eliminates poorly aligned positions and divergent regions of an alignment of DNA or protein sequences	0.91b	https://www.biologiaevolutiva.org/jcastresana/Gblocks.html		
MAFFT	MAFFT is a multiple sequence alignment program	7.526	https://mafft.cbrc.jp/alignment/software/		
muscle	Multiple Sequence Alignment faster and more accurate than clustalw	5.1	https://www.drive5.com/muscle/		
t_coffee	Multiple sequence alignment package	12.00.7	https://tcoffee.org/Projects/tcoffee/index.html		
minimap2	A versatile sequence alignment program that aligns DNA or mRNA sequences against a large reference database	2-2.30	https://github.com/lh3/minimap2		
	seqkit soap SRAtoolkit STAR Stringtie CaTupat CAT_pack ABySS ABySS AIIpaths-LG Canu Canu Canu Canu Canu Canu Canu Canu	SAM Tools provide various utilities for manipulating alignments in the SAM formatseqkitSecKit - a cross-platform and ultrafast toolkit for FASTA/Q file manipulationsoapShort Oligonuclectide Analysis PackageSRAtoolkitCalection of tools and libraries for using data in the INSDC Sequence Read Archives.STARSpliced Transcripts Alignment to a ReferenceStringtieStringtie is a fast and highly efficient assembler of RNA-Seq alignments into potential transcripts.TophatTopHat is a fast splice junction mapper for RNA-Seq readsAT_packA pipelines for the taxonomic classification of long DNA sequences and potential transcripts.ABySSAssembly By Short Sequences - a de novo, parallel, paired-end sequence assembler.Alipaths-LGThe new short read genome assembler.Alipaths-LGThe new short read genome assembler.AmanuelityMasureCA (Maryland Super-Read Celera Assembler) genome assembly toolfor Nanopore MinION).hiffasmMaSURCA (Maryland Super-Read celera Assembler) genome assembly toolfor containing various assembly toolfor containing various assembly toolfor Nanopore long noisy reads.SPAdesSequence assembler for very short readssoap denovoShort Oligonuclectide Analysis Package genome assembler.MutherA tool for (1) aligning two DNA sequences row reads coring parameters automaticallyMUMmerSayte as system for ruery short readssoap denovoSostembler Grue Alignmentsoap denovoSource alignment for proteinsclustalMuttiple Sequence Alignment <td>samtoolsSAM Tools provide various utilities for manipulating alignments in the SAM iormal context of the SAM tools provide various utilities for manipulation1.19.2seqkitSeqkit - a cross-platform and utrafast colocition of tools and libraries for using data in the INSDC Sequence Read Archives.2.21SRAtoolkitThe SRA Toolkit and SDK from NCBI is a collection of tools and libraries for using data in the INSDC Sequence Read archives.3.0.0STARSpliced Transcripts Alignment to a potential transcripts.3.0.0TophatSpliced Transcripts Alignment too potential transcripts.3.0.0TophatTopHat is a fast splice junction mapper for RNA-Seq reads3.0.0Alipaths-LGApipelines for the taxonomic cassembled genomes (MAGS parallel, paired-end sequence assembler sequence assembler oxford Nanopore MinION).3.3.4Alipaths-LGThe new short read genome assembler sequence for high-noise single-molecule sequence seembler for Nanopore long noisy row frags3.1.1NEGATNeSURCA (Maryland Super-Read Celera sesembler for Nanopore long noisy row frags3.1.1SPAdesS.PAdes St. Petersburg genome assembly tool Na sequences assembly tool Na nopore long noisy row frags3.1.1Spade in Novel method for the efficient assembler for Na-sequences and (2) infer</td>	samtoolsSAM Tools provide various utilities for manipulating alignments in the SAM iormal context of the SAM tools provide various utilities for manipulation1.19.2seqkitSeqkit - a cross-platform and utrafast colocition of tools and libraries for using data in the INSDC Sequence Read Archives.2.21SRAtoolkitThe SRA Toolkit and SDK from NCBI is a collection of tools and libraries for using data in the INSDC Sequence Read archives.3.0.0STARSpliced Transcripts Alignment to a potential transcripts.3.0.0TophatSpliced Transcripts Alignment too potential transcripts.3.0.0TophatTopHat is a fast splice junction mapper for RNA-Seq reads3.0.0Alipaths-LGApipelines for the taxonomic cassembled genomes (MAGS parallel, paired-end sequence assembler sequence assembler oxford Nanopore MinION).3.3.4Alipaths-LGThe new short read genome assembler sequence for high-noise single-molecule sequence seembler for Nanopore long noisy row frags3.1.1NEGATNeSURCA (Maryland Super-Read Celera sesembler for Nanopore long noisy row frags3.1.1SPAdesS.PAdes St. Petersburg genome assembly tool Na sequences assembly tool Na nopore long noisy row frags3.1.1Spade in Novel method for the efficient assembler for Na-sequences and (2) infer		

Aligniysis Type	Application name	Description	installed ver.	Official URL			
	miniprot	Miniprot aligns a protein sequence against a genome with affine gap penalty, splicing and frameshift		https://github.com/lh3/miniprot			
Datbase search	dbget	DBGET is an integrated database retrieval system for major biological databases		https://www.kanehisa.jp/ja/about_dbget_ja.html			
Sequence Assembler	САРЗ	Multiple Sequence Alignment	122107	https://faculty.sites.iastate.edu/xqhuang/cap3-and-pcap- sequence-and-genome-assembly-programs			
	consed	Assembly Editor	29.0	http://bozeman.mbt.washington.edu/consed/consed.html			
	Phrap	Phrap is a program for assembling shotgun DNA sequence data	1.090518	http://www.phrap.org/phredphrapconsed.html			
	Phred	The phred software reads DNA sequencing trace files, calls bases, and assigns a quality value to each called base	071220	http://www.phrap.org/phredphrapconsed.html			
	TGICL	Multiple Sequence Alignment (for huge data set)	2.1	it's gone : http://compbio.dfci.harvard.edu/tgi/software/			
	Augustus	AUGUSTUS is a program that predicts genes in eukaryotic genomic sequences	3.5.0	https://github.com/Gaius-Augustus/Augustus			
	Genemark	A family of gene prediction programs	4.69	https://genemark.bme.gatech.edu/			
	genscan	Gene prediction		https://www.genes.mit.edu/GENSCAN.html			
	glimmer glimmerhmm	Glimmer is a system for finding genes in microbial DNA	30.02 3.0.4	https://ccb.jhu.edu/software/glimmerhmm/			
Gene prediction	TSEBRA	TSEBRA is a combiner tool that selects transcripts from gene predictions based on the support by extrisic evidence in form of introns and start/stop codons		https://github.com/Gaius-Augustus/TSEBRA			
	Metaeuk	MetaEuk - sensitive, high-throughput gene discovery and annotation for large-scale eukaryotic metagenomics	7-bba0d80	https://github.com/soedinglab/metaeuk			
	Prodigal	Prodigal: Fast, reliable protein-coding gene prediction for prokaryotic genomes.	2.6.3	https://github.com/hyattpd/Prodigal			
	HMMER	Biosequence analysis using profile HMM	3.4	http://hmmer.org/			
Motif search	Interproscan	A tool that combines different protein signature recognition methods into one resource	5.73-104.0	https://www.ebi.ac.uk/interpro/			
	meme	Multiple Em for Motif Elicitation	5.4.1	https://meme-suite.org/meme/index.html			
Functional annotation	eggNOG-Mapper	A tool for fast functional annotation of novel sequences.	2.1.12	https://github.com/eggnogdb/eggnog-mapper			
	mrbayes	MrBayes is a program for Bayesian inference and model choice across a wide range of phylogenetic and evolutionary models	3.2.7 3.2.7.mpi	https://nbisweden.github.io/MrBayes/			
	njplot	NJplot is a tree drawing program	2.4	https://doua.prabi.fr/software/njplot			
	paup	Tools for inferring and interpreting phylogenetic trees	4b10	https://paup.phylosolutions.com/			
phylogenetic tree analysis	Phylip	A package of programs for inferring phylogenies (evolutionary trees)	3.697	https://phylipweb.github.io/phylip/			
	PhyML	PhyML is a phylogeny software based on the maximum-likelihood principle	3.3	https://github.com/stephaneguindon/phyml			
	RAxML (raxmIHPC)	RAxML - Randomized Axelerated Maximum Likelihood	8.2.13	https://github.com/stamatak/standard-RAxML/			
	tree-puzzle	Program to reconstruct phylogenetic trees from molecular sequence data by maximum likelihood	5.3.rc16	http://www.tree-puzzle.de/			
Single cell analysis	CellRanger	A set of analysis pipelines that perform sample demultiplexing, barcode processing, single cell 3' and 5' gene counting, V(D)J transcript sequence assembly and annotation, and Feature Barcode analysis from single cell data.	8.0.1	https://www.10xgenomics.com/jp/support/software/cell- ranger/latest			
	MARVEL	MARVEL is an R package developed for alternative splicing analysis at single-cell resolution.	1.0.0	https://github.com/wenweixiong/MARVEL			
Repeat	RepeatMasker	RepeatMasker is a program that screens DNA sequences for interspersed repeats and low complexity DNA sequences	4.1.8	https://www.repeatmasker.org/			
	RepeatModeler	RepeatModeler is a de novo transposable element (TE) family identification and modeling package.	2.0.6	https://www.repeatmasker.org/RepeatModeler/			

Analysis Typ	e Application name	Description	installed ver.	Official URL
tRNA search	tRNAscan-SE	tRNAscan-SE pioneers the large-scale use of covariance models to annotate tRNA genes in genomes	2.0.5	https://trna.ucsc.edu/tRNAscan-SE/
Other	TransDecoder	TransDecoder identifies candidate coding regions within transcript sequences.	5.7.1	https://github.com/TransDecoder/TransDecoder

■ apptainer で利用するためのコンテナ.sif ファイルが用意されているもの

- 作業ディレクトリにシンボリックリンクを作ってご利用ください
- パス:

/apl/bio/container/

		Version	.sif file	remarks	Official URL
AGAT	AGAT has the power to check, fix, pad missing information (features/attributes) of any kind of GTF and GFF to create complete, sorted and standardised gff3 format.	1.4.1	AGAT/1.4.1/agat_1.4.1pl5321hdfd78af_0.sif		https://github.com/NBISweden/AGAT
BRAKER	BRAKER is a program that predicts genes in eukaryotic genomic sequences	3.0.2	BRAKER/3.0.2/braker3.sif		https://github.com/Gaius-Augustus/BRAKER
BUSCO	Assessing genome assembly and annotation completeness with single-copy orthologs	5.8.0	BUSCO/5.8.0/busco580.sif	Required Options : offline	https://busco.ezlab.org/
DeepConsensus	DeepConsensus uses gap-aware sequence transformers to correct errors in Pacific Biosciences (PacBio) Circular Consensus Sequencing (CCS) data.	1.2.0	DeepConsensus/1.2.0/deepconsensus.sif		https://github.com/google/deepconsensus
DeepTMHMM	DeepTMHMM is currently the most complete and best- performing method for the prediction of the topology of both alpha-helical and beta-barrel transmembrane proteins.	1.0.42	DeepTMHMM/1.0.42/deeptmhmm_edit_g.sif		https://dtu.biolib.com/DeepTMHMM
EpiTyping	EpiTyping is a tool for detecting imprinting and X- chromosome inactivation status from RNA-seq	1	EpiTyping/epityping.sif		https://github.com/Gal-Keshet/EpiTyping
GALBA	GALBA uses the protein sequences of several (few) or one closely related species to generate a training gene set for AUGUSTUS with either miniprot or GenomeThreader.	1.0.7	GALBA/1.0.7/galba107_aug35.sif	with Augustus 3.5.0	https://github.com/Gaius-Augustus/GALBA
GATK	The GATK is the industry standard for identifying SNPs and indels in germline DNA and RNAseq data.	4.0.1	GATK/4.0.1/gatk-4.sif		https://github.com/broadinstitute/gatk

ipyrad	An interactive assembly and analysis toolkit for restriction-site associated DNA (RAD-seq) and related data types.	0.9.81	ipyrad/0.9.81/ipyrad_0.9.81pyh5e36f6f_0	https://ipyrad.readthedocs.io/en/master/
PASA Pipeline	PASA, acronym for Program to Assemble Spliced Alignments (and pronounced 'pass- uh'), is a eukaryotic genome annotation tool that exploits spliced alignments of expressed transcript sequences to automatically model gene structures		/apl/bio/container/PASA/2.5.3/pasapipeline.v2.5.3.simg	https://github.com/PASApipeline/PASApipeline