

The 2018 *Nucleic Acids Research* database issue and the online molecular biology database collection

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ABSTRACT

The 2018 *Nucleic Acids Research* Database Issue contains 181 papers spanning molecular biology. Among them, 82 are new and 84 are updates describing resources that appeared in the Issue previously. The remaining 15 cover databases most recently published elsewhere. Databases in the area of nucleic acids include 3DIV for visualisation of data on genome 3D structure and RNArchitecture, a hierarchical classification of RNA families. Protein databases include the established SMART, ELM and MEROPS while GPCrdb and the newcomer STCRDab cover families of biomedical interest. In the area of metabolism, HMDB and Reactome both report new features while PULDB appears in NAR for the first time. This issue also contains reports on genomics resources including Ensembl, the UCSC Genome Browser and ENCODE. Update papers from the IUPHAR/BPS Guide to Pharmacology and DrugBank are highlights of the drug and drug target section while a number of proteomics databases including proteomicsDB are also covered. The entire Database Issue is freely available online on the *Nucleic Acids Research* website (<https://academic.oup.com/nar>). The NAR online Molecular Biology Database Collection has been updated, reviewing 138 entries, adding 88 new resources and eliminating 47 discontinued URLs, bringing the current total to 1737 databases. It is available at <http://www.oxfordjournals.org/nar/database/c/>.

NEW AND UPDATED DATABASES

This 2018 *Nucleic Acids Research* Database Issue is the 25th annual collection of bioinformatic databases. The quarter century arrives with 181 papers which, as ever, span all areas of molecular biology research. The total includes 82 new databases (Table 1) and 84 updates of resources that have previously appeared in the Database Issue. There are also 15

updates on databases previously described elsewhere (Table 2).

As in previous years, databases are grouped into eight broad subject categories. These cover (i) nucleic acid sequence and structure, transcriptional regulation; (ii) protein sequence and structure; (iii) metabolic and signalling pathways, enzymes and networks; (iv) genomics of viruses, bacteria, protozoa and fungi; (v) genomics of human and model organisms plus comparative genomics; (vi) human genomic variation, diseases and drugs; (vii) plants and (viii) other topics, such as proteomics databases. In an era of increasingly interdisciplinary research, it is no surprise that the content of many databases spans multiple categories so that resources often do not sit comfortably in a single category. Readers are again urged to browse the whole issue, rather than confining themselves to the most obviously relevant sections. The *Nucleic Acids Research* online Molecular Biology Database Collection, which is available at <http://www.oxfordjournals.org/nar/database/c/>, retains its more finely grained organisation, encompassing 15 categories and 41 subcategories.

The issue begins with broad surveys of resources at major global centres, including the U.S. National Center for Biotechnology Information (NCBI), the European Bioinformatics Institute (EBI) and the BIG Data Center at the Beijing Institute of Genomics, Chinese Academy of Sciences. The NCBI Resources paper (1) presents an interesting analysis illustrating the extent of the cross-talk between different databases within the site, exemplifying the value to the user of the extensive data integration implemented at these centres. The EBI paper (2) describes new data types including image data, biobanks and biosamples, as well as charting the continued exponential growth in the volume of many kinds of data. The newest of the three, the BIG Data Center (3), focuses on genomic information, but also hosts facilities for samples, program code, and wikis. Many of the wikis are very active and have previously featured in NAR eg lncRNAWiki (4).

The 'Nucleic acid databases' section begins with updates from the International Nucleotide Sequence Database Collaboration (5) and its three contributors, GenBank, ENA and DDBJ (6–8) which together face the challenge of con-

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Table 1. Descriptions of new online databases in the 2018 NAR Database issue

Database	URL	Brief description ^a
3DIV	http://kobic.kr/3div	3D-genome Interaction Viewer and database
AAgMarker	http://bioinfo.wilmer.jhu.edu/AAgMarker/index.jsp	Serum autoantigen biomarkers from proteome microarrays
aBiofilm	http://bioinfo.imtech.res.in/manojk/abiofilm/	Anti-biofilm compounds
ActiveDriverDB	https://activedriverdb.org/	Genome variation mapped against post-translational modifications
ADReCS-Target	http://bioinf.xmu.edu.cn/ADReCS-Target	Adverse Drug Reactions linked to proteins, genes and genetic variants
AmyPro	http://amypro.net	Proteins with validated amyloidogenic regions
anti-CRISPRdb	http://cefg.uestc.edu.cn/anti-CRISPRdb/	anti-CRISPR proteins
AraGWAS Catalog	https://aragwas.1001genomes.org	Arabidopsis Genome-Wide Association Studies
ASpedia	http://combio.snu.ac.kr/aspedia	Alternative Splicing Encyclopedia
ChannelsDB	http://ncbr.muni.cz/ChannelsDB	Channels, pores and tunnels found in biomacromolecular structures
CirGRDB	http://cirgrdb.biols.ac.cn	Regulation of RNAs in circadian rhythms
ClusterCAD	https://clustercad.jbei.org and http://clustercad.igb.uci.edu	Engineering of type I modular polyketide synthases
CR2Cancer	http://cis.hku.hk/CR2Cancer	Chromatin Regulators and Cancer
CSCD	http://gb.whu.edu.cn/CSCD	Cancer-Specific cRNA Database
dbCAN-seq	http://cys.bios.niu.edu/dbCAN-seq	Genome scale CAZymes and CAZyme gene clusters
dbCoRC	http://dbcorc.cam-su.org/	Core transcriptional Regulatory Circuit models
DifferentialNET	http://netbio.bgu.ac.il/diffnet	Differential protein-protein interactions in human tissues
DiseaseEnhancer	http://bioc.hrbmu.edu.cn/DiseaseEnhancer/	Enhancer-disease associations
DISNOR	http://disnor.uniroma2.it/	Protein interaction networks linking disease genes
DreamBASE	http://rna.sysu.edu.cn/dreamBase	Human expressed pseudogenes: DNA Modification, RNA Regulation and bound proteins
ECODrug	http://www.ecodrug.org	Evolutionary Conservation of Drug targets
EpiDenovo	http://61.148.58.210:8080/EpiDenovo/	The epigenome in mammalian embryonic development
EPD	https://peptracker.com/epd/	Encyclopedia of Protein Dynamics
EVLncRNAs	http://biophy.dzu.edu.cn/EVLncRNAs/	Experimentally Validated lncRNAs including disease indications
eRAM	http://www.unimd.org/eram/	Annotated rare diseases
ExoRBase	http://www.exoRBase.org	Human blood exosome RNAs
FlavorDB	http://cosylab.iitd.edu.in/flavordb	Flavour molecules
FusionDB	http://services.bromberglab.org/fusiondb/	Functional-repertoire similarity-based organism network
GVM	http://bigd.big.ac.cn/gvm/	Genome Variation Map
HCMDB	http://hcmdb.i-sanger.com/index	Human Cancer Metastasis DataBase
HEDD	http://zdzlab.einstein.yu.edu/1/hedd.php	Human Enhancer Disease Database
ICG	http://icg.big.ac.cn	Internal Control Genes for RT-qPCR normalization
IMOTA	https://ccb-web.cs.uni-saarland.de/imota/	Interactive Multi-Omics-Tissue Atlas
iPTMnet	http://research.bioinformatics.udel.edu/iptmnet/	Post-Translational Modification networks
ITSoneDB	http://itsonedb.cloud.ba.infn.it/	Eukaryotic ribosomal RNA Internal Transcribed Spacer 1 sequences
jMorp	https://jmorp.megabank.tohoku.ac.jp/	Metabolomics and proteomics of 1000 healthy Japanese people
LINCS Data Portal	http://lincsportal.ccs.miami.edu/dcic-portal/	Cell-based perturbation-response signatures
LinkedOmics	http://www.linkedomics.org	Multi-omics analysis of 32 cancers
Lnc2Meth	http://www.bio-bigdata.com/Lnc2Meth	lncRNAs and DNA methylation
m6AVar	http://m6avar.renlab.org/	Human variants affecting m6A sites
MeDReaders	http://medreader.org/	Transcription factors binding methylated DNA
microbiomeDB	http://microbiomeDB.org	Mining and analysing microbiome data
MINTbase	https://cm.jefferson.edu/MINTbase/	Mitochondrial and nuclear tRNA fragments
miRCarta	https://mircarta.cs.uni-saarland.de/	miRNAs and precursors
mirTrans	http://mcube.nju.edu.cn/jwang/lab/soft/mirtrans/	Cell-specific transcriptional information for human miRNAs
MIST	http://fgrtools.hms.harvard.edu/ProteinSearch/	Model organism molecular interaction data
MGA	http://cvg.vital-it.ch/mga/	Mass Genome Annotation
MMP	https://mmp.sfb.uit.no/databases/	Marine Metagenomics Portal
MSDD	http://www.bio-bigdata.com/msdd/	miRNA SNP Disease Database
mSignatureDB	http://tardis.cgu.edu.tw/msignaturedb	Mutational signatures in human cancers
MVP	http://mvp.medgenius.info	Microbe-phage interactions
NPASS	http://bidd2.nus.edu.sg/NPASS/	Natural Product quantitative Activities
OverGeneDB	http://overgenedb.amu.edu.pl	Overlapping protein-coding genes
PAMDB	http://pseudomonas.umaryland.edu	<i>Pseudomonas aeruginosa</i> Metabolome DataBase
PancanQTL	http://bioinfo.life.hust.edu.cn/PancanQTL/	Expression quantitative loci (eQTL) analysis of cancer samples
PedAM	http://www.unimd.org/pedam/	Pediatric Disease Annotation & Medicine
PCSD	http://systemsbiology.cau.edu.cn/chromstates	Plant Chromatin State Database
PGG.Population	https://www.pggpopulation.org	Genomic diversity of diverse human populations
PharmacODB	http://pharmacodb.pmgenomics.ca	Pharmacogenomics of cancer cell lines
PICKLES	http://pickles.hart-lab.org	Pooled In-vitro Crispr Knockout Library Essentiality Screens
PIT-DB	http://pitdb.org	Proteomics Informed by Transcriptomics
Planteome	http://www.planteome.org	Portal for plant ontologies and annotations
PopHuman	http://pophuman.uab.cat	Population genomics-oriented genome browser

Table 1. Continued

Database	URL	Brief description ^a
qPrimerDB	http://biodb.swu.edu.cn/qprimerdb	qPCR primers for 200 organisms
RISE	http://rise.zhanglab.net	RNA-RNA interactions
RNArchitecture	http://iimcb.genesilico.pl/RNArchitecture/	Structural classification of RNAs
SBCCDB	http://sbccdb.moffitt.org	Sleeping Beauty Cancer Driver DataBase
SCPortalen	http://single-cell.clst.riken.jp/	Human and mouse single-cell centric database
SEECancer	http://biocc.hrbmu.edu.cn/SEECancer	Evolutionary-stage specific somatic events in cancer
StemMapper	http://stemmapper.sysbiolab.eu	Stem cell gene expression
STCRDab	http://opig.stats.ox.ac.uk/webapps/stcrdab	Structural T-Cell Receptor Database
SystemMHC Atlas	https://systemhcatlas.org/	Immunopeptidomics of MHC-bound peptides
Tabloid Proteome	http://iomics.ugent.be/tabloidproteome	Protein associations inferred from Mass Spectrometry
Target-Pathogen	http://target.sbg.qb.fcen.uba.ar/patho	Drug target optimisation in pathogens
TC3A	http://tc3a.org/	3' untranslated regions, alternative polyadenylation and cancer
TissGDB	http://zhaobioinfo.org/TissGDB	Tissue-specific Gene DataBase in cancer
TranslatomeDB	http://www.translatomedb.net/	Translatome data from RNC-Seq and Ribo-Seq
TriForC database	http://bioinformatics.psb.ugent.be/triforc/	Triterpene pathways
TCSBN	http://inetmodels.com	Tissue and Cancer-Specific Biological Networks
VarCards	http://varcards.biols.ac.cn/	Interpretation of coding variants in the human genome
VDJdb	https://vdjdb.cdr3.net/	T-cell receptor sequences with known antigen specificity
Virus Taxonomy	http://ictv.global	Taxonomy of viruses

^aFor full references to the databases featured in this issue, please see the Table of Contents.

Table 2. Updated descriptions of databases most recently published elsewhere

Database	URL	Brief description ^a
BioMuta and BioExpress	https://hive.biochemistry.gwu.edu/biomuta and https://hive.biochemistry.gwu.edu/bioexpress	Cancer SNVs and gene expression
BioStudies	https://www.ebi.ac.uk/biostudies/	Data of all kinds relating to a single study
iSyTE	http://research.bioinformatics.udel.edu/iSyTE	Integrated Systems Tool for Eye gene discovery
miRandola	http://mirandola.iit.cnr.it/	Extracellular and circulating non-coding RNAs
mirDIP	http://ophid.utoronto.ca/mirDIP/	microRNA Data Integration Portal
MNDR	http://www.rna-society.org/mndr/	Mammal ncRNA-Disease Repository
NLSdb	https://roslab.org/services/nlsdb/	Nuclear Localization Signals
PAGER 2.0	http://discovery.informatics.uab.edu/PAGER/	Pathway, Annotated-list, and Gene-signature Electronic Repository
ProteomicsDB	https://www.ProteomicsDB.org	Mass spectrometry of the human proteome
PULDB	http://www.cazy.org/PULDB_new/	Polysaccharide Utilization Loci in Bacteroidetes species
ReMap	http://remap.cisreg.eu	Transcription factor ChIP-seq data
RMDB	http://rmdb.stanford.edu	Structure mapping of RNA
SuperDrug2	http://cheminfo.charite.de/superdrug2	Approved drugs
TRRUST v2	http://www.grnpedia.org/trrust/	Transcriptional regulatory interactions in human and mouse.
TumorFusions	http://www.tumorfusions.org	Tumour fusion genes

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continued exponential growth in nucleic acid sequence data. Transcription factors (TF) and transcriptional regulation are represented by a number of databases. The popular returning database of TF binding profiles, JASPAR (9), is published back to back with the ReMAP database (10) of TF ChIP-seq data: data from ReMAP contributed directly to JASPAR's improved coverage. With recent intense interest in the role of 3D chromatin structure in gene regulation, the 3DIV resource (11) for 3D genome interaction visualisation is timely. The key RNA database Rfam (12) contributes an update describing a move to content based on a set of reference genomes. Mirroring changes made in Pfam (13), this eliminates much unhelpful redundancy and allows for clearer taxonomic comparisons.

miRNA biology is strongly represented by updates from established databases such as DIANA-TarBase (14) and mirDIP (15), as well as new databases such as miR-Carda (16). The new MSDD (17) links miRNA SNPs to diseases while EVLncRNAs (18) and MNDR (19) also

major on disease links for non-coding RNAs. The well-established MODOMICS database of RNA modifications (20) is the subject of an update paper which, among other developments, reports on the availability of liquid chromatography/mass spectrometry data for modified nucleosides, facilitating profiling of such modifications by these methods. RNA structure is covered by the returning RMDB database (21), containing chemical mapping information that can be used to predict RNA secondary and tertiary structure, and the new RNArchitecture (22) which introduces a hierarchical organisation of RNA families with a focus on 3D structures, in the manner popularised by protein databases like SCOP.

In the section on protein sequence and structure databases, the venerable SMART database celebrates 20 years with an update paper (23). It describes a particularly valuable new visualisation option, whereby domain architecture information can be added to phylogenetic trees with the Interactive Tree of Life (iTOL) tool (24). Another up-

date from PDBe (25) includes mention of a newly developed library of freely available web components for interactive data visualizations. One of these, the LiteMol 3D viewer, notably allows convenient display of electron density in the browser window. An update on the popular ELM database of protein sequence motifs (26) reports, among other developments, on how fascinating examples of bacterial pathogen mimicry of eukaryotic motifs are now included in the database. A new arrival, ChannelsDB (27) contributes our cover image and describes the channels, tunnels and pores in protein structures that allow substrate access to buried catalytic sites, for example, or molecular passage through a transmembrane protein. Certain protein classes or families justify their own bespoke databases through medical or biological importance. T-cell receptors are served in this issue by both VDJdb (28), focussing on receptor sequences of known specificity, and STCRDab (29) which collects and curates structural information, linking to and allowing searches against a wide variety of structural, sequence and functional data. The returning database GPCRdb (30), for G protein-coupled receptors, majors on carefully made homology models and mapping receptors to ligands.

Important updates in the metabolic and signalling section include the human metabolomics database HMDB (31). Release 4.0 brings huge increases in content, an improved interface and new kinds of information—predicted mass spectra and pharmacometabolomics. This issue also reports on a new metabolomics database, PAMDB (32), devoted to the bacterial pathogen *Pseudomonas aeruginosa*, justified not only by the biomedical importance of the organism but also by the novel metabolites that it contains. Metabolic pathways are covered by the well-known returning databases Reactome (33) and WikiPathways (34). The former update is notable for its Enhanced High Level Diagrams which superbly contextualise low-level pathways using images of cells, tissues and organs. Among enzyme-oriented databases MEROPS (35), devoted to proteases and their inhibitors, makes a welcome return with a near-doubling of sequences and cross-references to the PANTHER database (36). PANTHER full-length sequence based clustering is shown to be complementary to MEROP's domain-based structure. Carbohydrate-active enzymes are covered by the arrival in NAR of PULDB (37), covering polysaccharide utilization loci in the prominent gut bacteria of the phylum Bacteroidetes, and dbCAN-seq (38), which usefully extrapolates information from the well-known CAZy database (39) to a genome scale. At the enzyme mechanism level, this issue sees the merger of two databases, MACiE and CSA, each veterans of multiple Database Issues, into a single new resource M-CSA (Mechanism and Catalytic Site Atlas) (40).

In the microbial genomics section, there is an update paper from the yeast-focused SGD (41) which now includes curated lists of yeast genes that can replace the functions of human counterparts or vice versa. The popular TADB, covering toxins and antitoxins, also presents an update (42), as does *SubtiWiki* (43), devoted to the biology of *Bacillus subtilis*. Two new databases address viruses. The Virus Taxonomy (44) appears in NAR for the first time, despite the International Committee behind it dating back to the 1960s.

The second, MVP (45) describes the complex interactions between microbes and the phage clusters that can infect one or more of them.

Human and model organism genomics are strongly represented. The core resources Ensembl (46) and the UCSC Genome Browser (47) present their usual updates. The former is supplemented by an Ensembl Genomes paper (48) covering non-vertebrates which reports ~20 000 new genomes covered. Other well-known returning databases include ENCODE (49), RefSeq (50) and Genomicus (51), the last showcasing new karyotype evolutionary trees. Among new databases, current trends in cell and molecular biology are reflected in StemMapper (52) that focusses specifically on stem cell gene expression, and SCPortalen (53) which stores transcriptomics data, metadata and cell images at the single cell level. Another notable new arrival is PICKLES (54) which collects information on human gene essentiality from the results of genome scale CRISPR knockout and shRNA knockdown experiments in cancer and other cell lines.

As ever, databases devoted to human genomic variation and biomedical research are very well represented. Important returning databases include the IUPHAR/BPS Guide to Pharmacology (55) which covers properties of existing and potential drug targets. The authors of the update also describe a major new sister resource, the Guide to Immunopharmacology. An interesting evolutionary perspective on drug targets is provided by ECOdrug (56) which maps the presence or absence of drug target orthologues across species. This will help in efforts to address ecotoxicology concerns over binding of drugs to non-target wild species and assist with appropriate species choices for ecological risk assessments. The popular DrugBank (57) also returns, now in release 5.0 and bringing huge increases in data volume, new data types such as pharmacotranscriptomics and content reporting on the status of clinical trials. A major new resource is the Genome Variation Map (58) from the BIG Data Center covering 19 species. Its arrival is particularly timely with the announcement that comparable NCBI resources dbSNP and dbVar are to stop accepting non-human submissions (<https://ncbiinsights.ncbi.nlm.nih.gov/2017/05/09/phasing-out-support-for-non-human-genome-organism-data-in-dbsnp-and-dbvar/>). The well-used ClinVar resource (59) also contributes an update and is joined in interpreting human genome variation and its implications for disease by the newcomer VarCards (60). Two interesting new databases, PGG.Population (61) and PopHuman (62) present a population genomics perspective of human genome variation, each containing thousands of human genomes from across the world and allowing interactive exploration of and comparison between populations.

Plant databases represented here include the comparative genomics resources PLAZA (63) and Gramene (64). A major new Arabidopsis resource arrives in the form of the AraGWAS catalog (65) which contains hundreds of thousands of links between SNPs and curated phenotypes. In the last section proteomics databases are well-represented. An update is presented on the major quantitative proteomics resource proteomicsDB (66). Its protein-centric view links to an impressive variety of visualisations and to different

kinds of omics data. Future plans include an extension from its current human focus to model organisms. An intuitive user interface is also a strong point of the new EPD database (67), while PIT-DB (68) explicitly works at the intersection of RNA-seq transcriptomics and proteomics mass spectrometry. After covering such a variety of biological areas, it seems appropriate to finish with mention of the BioStudies database (69) that collates data of any and all kinds relating to a single study.

NAR ONLINE MOLECULAR BIOLOGY DATABASE COLLECTION

We reach this year the 25th update of the NAR online Molecular Biology Database Collection (which is freely available at <http://www.oxfordjournals.org/nar/database/c/>), featuring 88 new databases (Table 1) and 15 databases not described previously in the NAR Database Issue (Table 2). Within our ongoing verification processes to make sure information is still relevant, we have removed 47 obsolete or discontinued databases. After contacting their authors, 138 database entries have been updated with respect to new URLs, new descriptions, and/or other metadata.

We welcome suggestions for inclusion in the Collection of additional databases that have been published in other journals. Such suggestions should be addressed to XMF at xose.m.fernandez@gmail.com and should include database summaries in plain text, organized in accordance with the <http://www.oxfordjournals.org/nar/database/summary/1> template.

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