Frank Oliver Glöckner

List of Publications by Year in descending order

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51 papers

38,338 citations

27 h-index

201674

223800 46 g-index

56 all docs

56 docs citations

56 times ranked 41957 citing authors

#	Article	IF	CITATIONS
1	Implementing FAIR data management within the German Network for Bioinformatics Infrastructure (de.NBI) exemplified by selected use cases. Briefings in Bioinformatics, 2021, 22, .	6.5	18
2	The ELIXIR Core Data Resources: fundamental infrastructure for the life sciences. Bioinformatics, 2020, 36, 2636-2642.	4.1	47
3	Genomic characterization of filamentous phage vB_Vpal_VP-3218, an inducible prophage of Vibrio parahaemolyticus. Marine Genomics, 2020, 53, 100767.	1.1	10
4	Fast and accurate average genome size and 16S rRNA gene average copy number computation in metagenomic data. BMC Bioinformatics, 2019, 20, 453.	2.6	15
5	On-Site Analysis of Bacterial Communities of the Ultraoligotrophic South Pacific Gyre. Applied and Environmental Microbiology, 2019, 85, .	3.1	27
6	Meta-omics data and collection objects (MOD-CO): a conceptual schema and data model for processing sample data in meta-omics research. Database: the Journal of Biological Databases and Curation, 2019, 2019, .	3.0	14
7	The UNITE database for molecular identification of fungi: handling dark taxa and parallel taxonomic classifications. Nucleic Acids Research, 2019, 47, D259-D264.	14.5	2,072
8	The de.NBI / ELIXIR-DE training platform - Bioinformatics training in Germany and across Europe within ELIXIR. F1000Research, 2019, 8, 1877.	1.6	1
9	The de.NBI / ELIXIR-DE training platform - Bioinformatics training in Germany and across Europe within ELIXIR. F1000Research, 2019, 8, 1877.	1.6	3
10	A comprehensive fungi-specific 18S rRNA gene sequence primer toolkit suited for diverse research issues and sequencing platforms. BMC Microbiology, 2018, 18, 190.	3.3	84
11	The North Sea goes viral: Occurrence and distribution of North Sea bacteriophages. Marine Genomics, 2018, 41, 31-41.	1.1	17
12	Taxonomic and functional diversity of a coastal planktonic bacterial community in a river-influenced marine area. Marine Genomics, 2017, 32, 61-69.	1.1	15
13	A phylogenetic framework for the kingdom Fungi based on 18S rRNA gene sequences. Marine Genomics, 2017, 36, 33-39.	1.1	47
14	<i>UniEuk</i> : Time to Speak a Common Language in Protistology!. Journal of Eukaryotic Microbiology, 2017, 64, 407-411.	1.7	74
15	25 years of serving the community with ribosomal RNA gene reference databases and tools. Journal of Biotechnology, 2017, 261, 169-176.	3.8	679
16	Recurring patterns in bacterioplankton dynamics during coastal spring algae blooms. ELife, 2016, 5, e11888.	6.0	414
17	<i>Candidatus</i> Desulfofervidus auxilii, a hydrogenotrophic sulfateâ€reducing bacterium involved in the thermophilic anaerobic oxidation of methane. Environmental Microbiology, 2016, 18, 3073-3091.	3.8	115
18	Phylogeny-aware identification and correction of taxonomically mislabeled sequences. Nucleic Acids Research, 2016, 44, 5022-5033.	14.5	97

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19	Habitat and taxon as driving forces of carbohydrate catabolism in marine heterotrophic bacteria: example of the model algaeâ€associated bacterium ⟨i>Zobellia galactanivorans⟨ i> Dsij⟨sup>T⟨ sup⟩. Environmental Microbiology, 2016, 18, 4610-4627.	3.8	131
20	MyOSD 2014: Evaluating Oceanographic Measurements Contributed by Citizen Scientists in Support of Ocean Sampling Day. Journal of Microbiology and Biology Education, 2016, 17, 163-171.	1.0	6
21	JSpeciesWS: a web server for prokaryotic species circumscription based on pairwise genome comparison. Bioinformatics, 2016, 32, 929-931.	4.1	2,023
22	Identification of Habitat-Specific Biomes of Aquatic Fungal Communities Using a Comprehensive Nearly Full-Length 18S rRNA Dataset Enriched with Contextual Data. PLoS ONE, 2015, 10, e0134377.	2.5	62
23	Metatranscriptome of marine bacterioplankton during winter time in the North Sea assessed by total RNA sequencing. Marine Genomics, 2015, 19, 45-46.	1.1	14
24	The ocean sampling day consortium. GigaScience, 2015, 4, 27.	6.4	185
25	Minimum Information about a Biosynthetic Gene cluster. Nature Chemical Biology, 2015, 11, 625-631.	8.0	715
26	Niches of two polysaccharide-degrading <i>Polaribacter</i> isolates from the North Sea during a spring diatom bloom. ISME Journal, 2015, 9, 1410-1422.	9.8	182
27	Genomic Encyclopedia of Bacteria and Archaea: Sequencing a Myriad of Type Strains. PLoS Biology, 2014, 12, e1001920.	5.6	190
28	The founding charter of the Genomic Observatories Network. GigaScience, 2014, 3, 2.	6.4	51
29	Permanent draft genome of Rhodopirellula rubra SWK7. Marine Genomics, 2014, 13, 11-12.	1.1	6
30	Permanent draft genome of Rhodopirellula sallentina SM41. Marine Genomics, 2014, 13, 17-18.	1.1	5
31	The SILVA and "All-species Living Tree Project (LTP)―taxonomic frameworks. Nucleic Acids Research, 2014, 42, D643-D648.	14.5	2,667
32	Diversity and activity of marine bacterioplankton during a diatom bloom in the North Sea assessed by total RNA and pyrotag sequencing. Marine Genomics, 2014, 18, 185-192.	1.1	84
33	Uniting the classification of cultured and uncultured bacteria and archaea using 16S rRNA gene sequences. Nature Reviews Microbiology, 2014, 12, 635-645.	28.6	2,000
34	FastaValidator: an open-source Java library to parse and validate FASTA formatted sequences. BMC Research Notes, 2014, 7, 365.	1.4	2
35	Permanent draft genomes of the Rhodopirellula maiorica strain SM1. Marine Genomics, 2014, 13, 19-20.	1.1	5
36	Permanent draft genomes of the two Rhodopirellula europaea strains 6C and SH398. Marine Genomics, 2014, 13, 15-16.	1.1	4

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37	Permanent draft genomes of the three Rhodopirellula baltica strains SH28, SWK14 and WH47. Marine Genomics, 2014, 13, 13-14.	1.1	3
38	Genomic Standards Consortium Projects. Standards in Genomic Sciences, 2014, 9, 599-601.	1.5	26
39	Genomics in marine monitoring: New opportunities for assessing marine health status. Marine Pollution Bulletin, 2013, 74, 19-31.	5.0	196
40	The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Research, 2012, 41, D590-D596.	14.5	21,425
41	SINA: Accurate high-throughput multiple sequence alignment of ribosomal RNA genes. Bioinformatics, 2012, 28, 1823-1829.	4.1	2,826
42	Minimum information about a marker gene sequence (MIMARKS) and minimum information about any (x) sequence (MIxS) specifications. Nature Biotechnology, 2011, 29, 415-420.	17.5	608
43	Ecogenomics and genome landscapes of marine Pseudoalteromonas phage H105/1. ISME Journal, 2011, 5, 107-121.	9.8	62
44	Marine microbial genomics in Europe: current status and perspectives. Microbial Biotechnology, 2010, 3, 523-530.	4.2	28
45	The All-Species Living Tree project: A 16S rRNA-based phylogenetic tree of all sequenced type strains. Systematic and Applied Microbiology, 2008, 31, 241-250.	2.8	884
46	Mg-Traits pipeline: advancing functional trait-based approaches in metagenomics. ARPHA Conference Abstracts, 0, 4, .	0.0	0
47	The SILVA Database Project: An ELIXIR core data resource for high-quality ribosomal RNA sequences. Biodiversity Information Science and Standards, 0, 3, .	0.0	11
48	FAIR data in meta-omics research: Using the MOD-CO schema to describe structural and operational elements of workflows from field to publication. Biodiversity Information Science and Standards, 0, 3, .	0.0	3
49	Mainstreaming Molecular Biodiversity: A call for a unified and interoperable framework. Biodiversity Information Science and Standards, 0, 3, .	0.0	0
50	NFDI4BioDiversity: Biodiversity, ecology and environmental data. Biodiversity Information Science and Standards, 0, 3, .	0.0	1
51	Recommendations for connecting molecular sequence and biodiversity research infrastructures through ELIXIR. F1000Research, 0, 10, 1238.	1.6	3