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MicroScope—an integrated resource for community expertise of gene functions and comparative analysis of microbial genomic and metabolic data

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Abstract

The overwhelming list of new bacterial genomes becoming available on a daily basis makes accurate genome annotation an essential step that ultimately determines the relevance of thousands of genomes stored in public databanks. The MicroScope platform (http://www.genoscope.cns.fr/agc/microscope) is an integrative resource that supports systematic and efficient revision of microbial genome annotation, data management and comparative analysis. Starting from the results of our syntactic, functional and relational annotation pipelines, MicroScope provides an integrated environment for the expert annotation and comparative analysis of prokaryotic genomes. It combines tools and graphical interfaces to analyze genomes and to perform

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the manual curation of gene function in a comparative genomics and metabolic context. In this article, we describe the freeof-charge MicroScope services for the annotation and analysis of microbial (meta)genomes, transcriptomic and re-sequencing data. Then, the functionalities of the platform are presented in a way providing practical guidance and help to the nonspecialists in bioinformatics. Newly integrated analysis tools (i.e. prediction of virulence and resistance genes in bacterial genomes) and original method recently developed (the pan-genome graph representation) are also described. Integrated environments such as MicroScope clearly contribute, through the user community, to help maintaining accurate resources.

Key words: microbial genome annotation system; gene function curation; comparative genomics; transcriptomics; variant detection; metabolic networks

Introduction

Large-scale genome sequencing and the increasingly massive use of high-throughput approaches produce a vast amount of new information that completely transforms our understanding of thousands of species. However, despite the development of powerful bioinformatics approaches, full interpretation of the content of these genomes remains a difficult task. To address this challenge, several integrated environments that combine and standardize information from a variety of sources and apply uniform (re-)annotation techniques have been developed (i.e. EnsemblGenomes [1], IMG [2], PATRIC [3]). In the context of the French National Sequencing Center (CEA/DRF/Genoscope), we have developed the MicroScope platform, which is a software environment for management, annotation, comparative analysis and visualization of microbial genomes. Published for the first time in 2006 [4], the platform has been under continuous development within the LABGEM group at CEA, and its capacities are now extensive [5-7]. MicroScope serves different used cases in bioinformatics:

- · It supports the integration of newly sequenced or already available prokaryotic genomes through the offer of a free-of-charge service to the scientific community [genome annotation, RNA sequencing (RNA-seq) and variant analyses].
- It performs computational inferences including prediction of metabolic pathways, prediction of resistome and virulome, which can be used for genome analysis.
- It provides tools for (comparative) analyses and visualization of prokaryotic genomes.
- It supports collaborative expert annotation processes through the use of specific curation tools and graphical interfaces.

The present article provides a comprehensive description of MicroScope from the point of view of the end users. We start with the major objectives for which the platform was designed, and we give an overview of the main categories of MicroScope users and projects. Then we explain how to submit data and interact with the MicroScope team, and how to explore the annotated data, use the various analysis tools and perform expert annotation of gene functions. Technical details on the architecture of the system are given in the last section of this review. Where possible, earlier publications that provide more details are referenced. We conclude by one of the ongoing work that lead to a promising representation of the pan-genome of thousands of prokaryotic genomes.

Who is using MicroScope and for what purposes?

In the era of high-throughput sequencing technologies, a vast majority of genome sequences receive only automatic annotation, mainly based on sequence similarity, that can give spurious results [8]. Indeed, manual expertise of gene functions is a

time-consuming and expensive process, but it undoubtedly adds great value to resources. In knowledge bases such as UniProtKB [9], curation efforts remain restricted to large and widespread protein families, and these resources cannot replace expert curations made by specialized biologists in community systems, such as SEED [10], IMG [2] and MicroScope. Our integrated platform supports systematic and efficient revision of microbial genome annotation, data management together with comparative genomics and metabolic analyses [4-7]. The resource provides data from completed and ongoing genome projects together with post-genomic experiments (i.e. transcriptomics; re-sequencing of evolved strains; mutant collections) allowing users to improve the understanding of gene functions.

In comparison with other similar systems, MicroScope enables curation in a rich comparative genomic context and is mainly focused on (re-)annotation projects, which are built in close collaboration with microbiologists working on reference species. Indeed, MicroScope was initially dedicated to the annotation and analysis of Acinetobacter baylyi APD1 [11] and to biologists who do not have the required computing infrastructure to perform efficient annotation and analyses of newly sequenced bacterial genomes. Our system rapidly became a 'service' free of charge to the scientific community at large. From <400 user accounts in 2006, MicroScope counts >3300 personal accounts at present time (Figure 1). The number of registered users has doubled since 2013, and the platform has even widened its international popularity with 64% of accounts outside France. Many international projects are conducted through the platform involving users from distant geographic areas [7]. Although authentication is not required to navigate in MicroScope, it allows users to annotate genes and save data on their personal session. On average per month, we count 360 active accounts (i.e. the user logged in at least once in the month) and 2200 authentications among \sim 1700 monthly unique visitors.

The platform has been used to perform a complete expert annotation of several reference species such as Escherichia coli [12], Bacillus subtilis 128 [13, 14] and Pseudomonas putida KT2440 [15]. In addition, important pathogens and environmental species have also been extensively curated. The MicroScope system is now also used for variant analysis of re-sequenced bacterial strains (for example, in the context of bacterial evolution experiments) and for the analysis of transcriptomic experiments using RNA-seq sequencing data [6, 7], and finally, the platform is also (and in some cases, exclusively) used for the set of analysis tools pertaining to microbial genomics and metabolism, which have been integrated and made available through the MicroScope Web interface (see next sections). Indeed, the MicroScope platform has been cited 690 times since 13 years.

As shown in Figure 1, although the number of MicroScope users having a personal account has increased significantly since 2011, the number of expert annotations made each year is clearly decreasing, reaching only 21600 in 2016 (we registered >100000

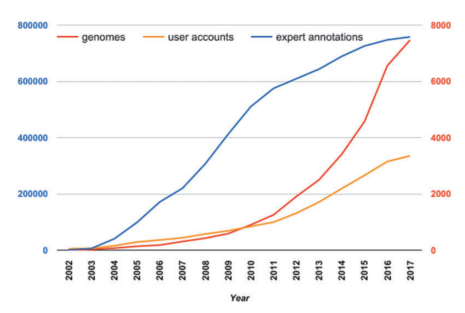


Figure 1. Evolution of the number of integrated genomes, user accounts and expert annotations stored in MicroScope since 2002. Red scale on the right refers to the number of integrated genomes (red curve) and to the number of user accounts (orange curve). Blue scale on the left refers to the cumulated number of expert annotations.

expert annotations in 2009). Past year, about one-tenth of the users performed curation of gene function and a third of them made >100 expert annotations. Obviously, with the number of prokaryotic genomes being sequenced today, the time-consuming task of expert annotation is totally unacceptable. This is the reason why our major efforts have been focused on the development of several key functionalities allowing to ease the expert annotation process and to notably improve the final annotation quality of the analyzed genomes, at least, for gene functions of interest.

An annotation service to researchers in microbiology

Interface for user data integration

Integration and analysis of genomic data into MicroScope are open and free of charge for the worldwide community of microbiologists. To standardize and make user submission fully automated, we have developed a dedicated Web interface (https://www.genoscope.cns.fr/agc/microscope/about/services. php). The service is mainly used for the annotation of microbial genomes: both newly sequenced genomes (which will remain private till the genome publication and/or their submission to public databanks) and, for comparative analysis purpose, public prokaryotic genomes (Figure 2). Moreover, three other types of services are provided for the integration of (i) genome assemblies (bins) from metagenomic samples (ii) RNA-seq data for quantitative transcriptomics and (iii) DNA sequencing (DNAseq) data to identify genomic variations in evolved strains (Figure 3). To ease data integration and comparative studies, standardization of contextual data about genome sequences is essential. For metagenomes, we have added a dedicated form that follows the MIMS specifications (minimum information about a metagenome sequence [16]). When submitting assembled metagenomic data in Microscope, the users are invited to select the type of environment (e.g. soil; air; water; human-associated; plant-associated) and to complete the associated fields (e.g. collection date, environment biome, geographic location, etc.). These fields are dynamically loaded and

displayed on metagenome type selection. Indeed, the MicroScope database model is flexible enough to store predefined descriptors, like MIMS, or the ones defined by users.

At present time, an average of eight genomes a day are requested for integration in the platform (this includes bins from metagenomic samples). The resource contains data for >7400 microbial genomes of which ~3100 are publicly available. In addition, 607 RNA-seq runs and 756 runs corresponding to the re-sequencing of evolved strains have also been requested for integration into MicroScope.

Running the annotation pipelines

About 25 analyses workflows include most of the currently used annotation software, plus some in-house tools and/or annotation strategies (Table 1). The newly sequenced (meta)genomes, generally submitted in several contigs and organized (or not) on the final chromosome(s), are first analyzed by the syntactic annotation pipeline to identify protein genes, transfer RNA (tRNA), ribosomal RNA (rRNA), noncoding RNA (ncRNA) and repeats (Figure 2, Table 1). For a more accurate prediction of small genes and/or atypical gene composition, we have developed a strategy to first construct appropriate gene models that takes into account the codon usage of the studied organism. These models are then used in the core of the AMIGene program [17]. Starting with the set of genomic objects identified during the syntactic annotation process, the next step is to infer biological functions of the predicted genes. Our functional annotation pipeline includes sequence similarity searches tools using generalist (i.e. UniProtKB/Swiss-Prot) or specialized (i.e. Interpro, FIGFAM, etc.) databases (Table 1). Results obtained with high-quality manually curated protein sequence data sets (i.e. Swiss-Prot, E. coli K-12, B. subtilis 168 MicroScope-curated genes) are first considered in the final functional automatic annotation procedure. This procedure also takes into account the results obtained from the computation of synteny groups with complete reference prokaryotic genomes and the one available in MicroScope. Indeed, for assigning function to novel proteins, gene context approaches often complement the classical homology-based

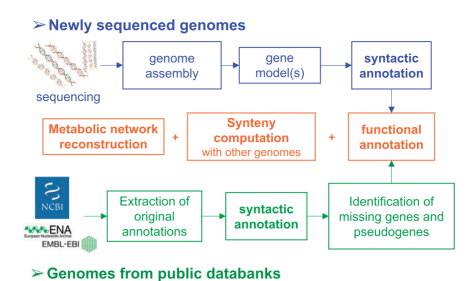


Figure 2. Annotation pipelines for the analysis of newly sequenced genomes and genomes already annotated in public databanks.

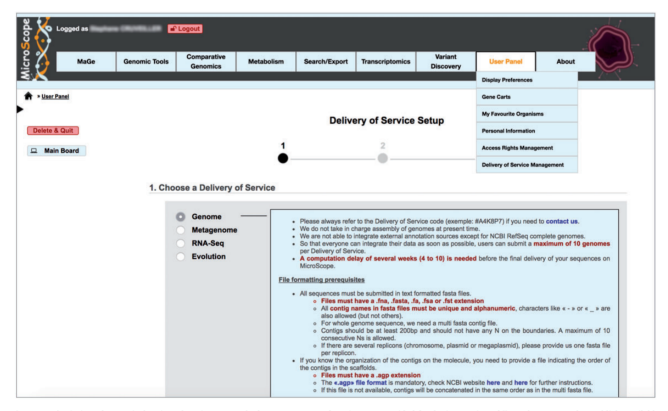


Figure 3. Submission of genomic data into the MicroScope platform. Four types of services are provided for the integration of (i) newly sequenced or publicly available genomes (Genome), (iii) genome assemblies/bins from metagenomic samples (Metagenome), (iiii) RNA-seq data for quantitative transcriptomics (RNA-Seq), (iv) DNAseq data to identify genomic variations in evolved strains (Evolution). Following the three main steps of the procedure, the user is invited to complete the requested metadata to describe sequencing, genomes and experimental properties, to upload FASTA (genome assemblies) or FASTQ (RNA-seq or DNA-seq reads) files and, finally, to approve the terms of services. Users are then informed by an e-mail about the progress of their integration request.

gene annotation in prokaryotes. The method we have developed offers the possibility of retaining more than one homologous gene (i.e. not only the bidirectional best hit), to allow for multiple correspondences between genes; that way, paralogy relations and/or gene fusions are easily detected [4].

Information from the syntactic and functional annotation pipelines can be placed into a biological context to understand how the predicted objects interact in functional modules such as metabolic pathways. Each genome integrated into MicroScope is processed by an in-house workflow based on the MetaCyc reference database [18] and on the Pathway Tools software [19]. This software creates a Pathway Genome DataBase (PGDB) containing the predicted pathways and reactions of an organism. It uses a matching procedure for which we directly use as input the official MetaCyc reaction frame identifiers when available in the genome annotation; this allows to avoid

Table 1. Software and databases integrated in the MicroScope pipelines

Topic	Name	Software	Database	Description	Internal	URL
Syntactic annotation	AMIGene Glimmer	x x		CoDing sequences (CDS) prediction	х	http://www.genoscope.cns. fr/agc/tools/amigene https://ccb.jhu.edu/soft
	Prodigal	x				ware/glimmer http://prodigal.ornl.gov
	MICheck	X		INSDC genome CDS re-	x	http://www.genoscope.cns.
				annotation		fr/agc/tools/micheck
	tRNAscan-SE	х		tRNA prediction		http://eddylab.org/soft ware/tRNAscan-SE
	RNAmmer	х		rRNA prediction		http://www.cbs.dtu.dk/ser vices/RNAmmer
	Rfam/Infernal	х	х	ncRNA families and prediction		http://rfam.xfam.org, http:// eddylab.org/infernal
	RepSeek	х		DNA sequence repeats		http://wwwabi.snv.jussieu. fr/public/RepSeek
	Alien hunter	х		DNA compositional biases to detect HGT regions		http://www.sanger.ac.uk/sci ence/tools/alien-hunter
	SIGI-HMM	х				http://www.brinkman.mbb. sfu.ca/~mlangill/sigi- hmm
	GenProtFeat	x		Gene/protein features	x	*******
	Taxonomy		х	NCBI taxonomy database		https://www.ncbi.nlm.nih. gov/taxonomy
Functional	BLAST+	X		DNA/protein sequence		https://blast.ncbi.nlm.nih.
annotation	Diamond	х		alignment		gov https://github.com/bbuch
	UniProtKB		x	Protein sequence and func-		fink/diamond http://www.uniprot.org
	InterPro	x	x	tion database Protein signature and fam-		https://www.ebi.ac.uk/
	COG	x	x	ily prediction Protein family annotation		interpro https://www.ncbi.nlm.nih.
	FigFam	х	х	and prediction		gov/COG http://www.nmpdr.org/FIG/
	MICFAM	x		Protein sequence family	x	wiki/view.cgi/FIG/FigFam
	WIIGI 7 IWI	A		classification with SiliX	A	
	SiliX	х		Clustering of protein sequences		https://lbbe.univ-lyon1.fr/- SiLiXhtml
	ENZYME		X	Enzymatic activity database		http://enzyme.expasy.org
	PRIAM	х		Enzymatic activity prediction		http://priam.prabi.fr
	dbCAN	Х		Carbohydrate-active en- zyme prediction		http://csbl.bmb.uga.edu/ dbCAN/
	SignalP	Х		Signal peptide cleavage site prediction		http://www.cbs.dtu.dk/ser vices/SignalP
	TMHMM	x		Transmembrane helix prediction		http://www.cbs.dtu.dk/ser vices/TMHMM
	LipoP	x		Lipoprotein prediction		http://www.cbs.dtu.dk/ser vices/LipoP
	PSORTb	x		Subcellular localization prediction		http://www.psort.org
	VFDB		X	Virulence factor database		http://www.mgc.ac.cn/VFs
	VirulenceFinder		х			https://cge.cbs.dtu.dk/ser vices/VirulenceFinder
	CARD/RGI	X	X	Antibiotic resistance data- base and prediction		https://card.mcmaster.ca
	AutoFassign	x		Automatic functional anno- tation of proteins	X	
Relational annotation	Syntonizer	x		Synteny conservation detection	х	http://www.inrialpes.fr/ helix/people/viari/ cccpart/

Table 1. Continued

Topic	Name	Software	Database	Description	Internal	URL
	Directon	х		Operon prediction	х	
	PhyloProfile	х		Phylogenetic profilef co- evolution score	Х	https://dx.doi.org/10.1186 %2F1471-2164-13-69
	RGP	x		Genomic plasticity region detection	Х	
	Pathway synteny	х		Synteny involved in meta- bolic pathways	Х	
	MIBiG/ antiSMASH	х	x	Biosynthetic Gene Cluster database and prediction		http://www.secondarymeta bolites.org/
	ChEBI		x	Chemical compound database		https://www.ebi.ac.uk/ chebi
	Rhea		х	Reaction database		http://www.rhea-db.org
	KEGG		x	Metabolic pathway database		http://www.genome.jp/kegg
	MetaCyc/ Pathway tools	х	х	Metabolic pathway data- base and prediction		https://metacyc.org, http:// brg.ai.sri.com/ptools/
Transcriptomics and variant	SSAHA2	х		Read mapping		http://www.sanger.ac.uk/sci ence/tools/ssaha2-0
discovery	BWA	х				https://github.com/lh3/bwa
	SAMtools	x		Mapping analysis		http://www.htslib.org/
	bedtools	х				http://bedtools.readthe docs.io
	PALOMA	x		Variant detection	х	
	DESeq	х		Differential gene expression analysis		http://bioconductor.org/ packages/release/bioc/ html/DESeq.html

overpredicted or missed enzymatic reactions [20]. The collection of MicroScope PGDBs is made available at the MicroCyc Web site (http://www.genoscope.cns.fr/agc/microcyc) and in the MicroScope database (see 'Exploration of metabolic data' section). Moreover, these metabolic networks are synchronized each night with new MicroScope genomes and expert annotations.

When a public prokaryotic genome is integrated into MicroScope, the original annotations are stored in the database, and the syntactic re-annotation process, which uses the MICheck procedure, often allows to identify missing genes or wrongly annotated one [21]. This step is useful to annotate more completely the pseudogenes found in a genome ('real' or because of sequencing errors), an important piece of information when comparing closely related species. Data from genomes available in public databanks generally remain with the 'public' status too in MicroScope.

A MicroScope staff to support and train a user community

As soon as annotations and comparative analysis results are processed by MicroScope, the user who submitted the genome(s) is alerted by an e-mail; he/she can subsequently use a specific administration tool to grant access to his/her collaborators and to define consultation and modification rights on the sequences ('User Panel' menu/'Access Rights Management' functionality). Continuing support and assistance to MicroScope users remain an important activity in the context of our services (or collaborative projects). These regular exchanges, together with the satisfaction surveys, are the most efficient way of performing continual evolution of the platform in response to user needs. Indeed, in addition to the user-

friendliness of the tools integrated into the platform (see below), the short response time and the quality of feedback to individual queries are highly appreciated aspects of the MicroScope service.

Microbiologists who submitted genomic data to the MicroScope platform are warmly invited to follow a training course organized by our team. Using the data related to their own project, attendees learn how to change or correct the current automatic functional annotations, and how to perform effective searches and analyses with the functionalities available through the Web interface. About twice a year, we provide for new users a four-and-a-half-day training 'Annotation and analysis of prokaryotic genomes using the MicroScope platform'. Since 2016, we also provide an advanced course for former trainees, so that they can remain up-to-date on recent developments. Since 2008, 450 users from 20 countries have been trained and 13 external sessions have been organized in France and abroad (Tunisia; Denmark; Germany; Switzerland; Spain; the Netherlands; China). More information is available on our Web site: http://www.genoscope.cns.fr/agc/microscope/training.

Data integration, service continuity and data conservation (backups) are currently provided free of charge. MicroScope services follow the quality management system of our laboratory (ISO 9001:2008 and NF X50-900:2013 standards).

All the data previously described (primarily genomes, analysis results and annotations) should be made appropriately accessible to biologist users, to allow efficient curation of annotations and to develop hypotheses about specific genomes or sets of genes to be experimentally tested. The following sections describe the MicroScope Web interface (http://www.genoscope.cns.fr/agc/microscope), i.e. the components accessible to our users, via secure or anonymous connections. For a complete description of each functionality in terms of input and output

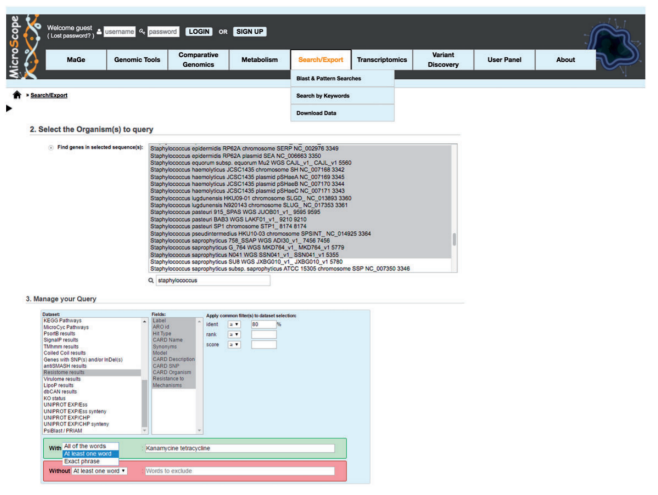


Figure 4. MicroScope interface illustrating the 'Search by keywords' functionality. In the 'multiple' mode, a set of Staphylococcus species has been selected, and the BLASTP similarity results obtained with well-known resistance genes stored in the CARD database are queried using an amino acid identity threshold of at least 80% and using the keywords 'kanamycine tetracycline'. The selection of 'At least one word' is required to apply an 'OR' between the two keywords.

data, a complete tutorial is available here: https://microscope. readthedocs.io.

Exploration of the genomic data: simple and advanced queries

The 'Search/Export' menu (Figure 4) allows the user to perform Blast and pattern searches in the MicroScope database, and to download, in standard file formats (Genbank, EMBL, GFF, etc.), sequences, annotation data and the metabolic networks. The 'Search by keywords' functionality allows the user to identify genes and functions of interest using a variety of selection filters. The 'single mode' is used to query only one chromosome and the 'multiple mode' to query several replicons (of one organism) and/or several genomes. A basic keyword search enables the user to quickly retrieve genes having a particular function (i.e. 'kinase', 'transporter'). Each kind of precomputed results (i.e. Blast results on various primary data, InterPro and FigFAM results, etc.) can be queried. Figure 4 shows an example of a query on the similarity searches in the CARD database [22] ('Resistome' data set).

Keyword searches are useful to compare current annotation of the gene functions with the results, in terms of biological function, given by a specific analysis method. Indeed, the result of a query can be refined with a further query. For example, one can search for gene annotated as 'protein of unknown function' (first query) and then, search for the one having significant Blast results with proteins annotated with specific functions (second query).

Whatever the query, the result output is a list of candidate genes, the genomic contexts of which can be easily visualized: next to the gene label, a magnify icon can be clicked to come back to the MaGe graphical representation with automatic displacement of the genome browser centered on the gene of interest.

MaGe (Magnifying Genome): a genome browser in the light of synteny results

The MaGe graphical interface is one of the functionality that had a strong positive resonance among users: this genome browser offers gene context exploration of the studied genome compared against other microbial genomes. The graphical representation of the synteny groups allows the user to quickly see if part of the genome being annotated shares similarities and locally conserved organization with the selected sequences. As shown in Figure 5, there is a clear synteny break in the visualized part of the E. coli CFT073 strain: the genes located between 5116000 and 5131000 share homologs only with the E. coli pathogenic strain ABU and, partially, with the E. coli commensal strain

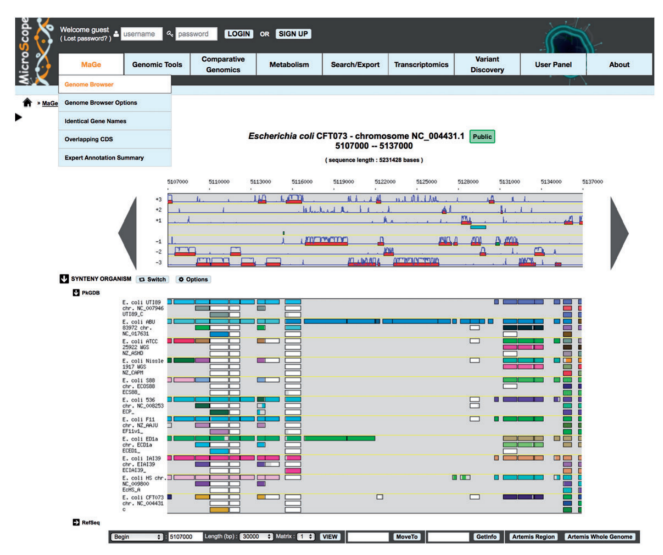


Figure 5. MicroScope genome browser and synteny map. The first graphical map contains part of the genome being analyzed (here 30 kb of E. coli CFT073), over which the user can navigate (moving and zooming functionalities). The predicted coding genes are drawn, on the six reading frames, in red rectangles together with the coding prediction curves (computed with the gene model selected by the user; 'Matrix' selection menu). Below this genome browser, is represented the synteny map in which each line shows the similarity results between the genome being annotated (E. coli CFT073) and other selected genomes (i.e. 11 pathogenic and commensal E. coli strains; the selection is performed using the 'Options' functionality). On this map, a rectangle flags the existence of a gene, somewhere in the compared genome, homolog to the corresponding gene in the genome browser. If, for several co-localized CDSs on the annotated genome, there are several co-localized homologs on the compared genome, the rectangles are all of the same color; otherwise, the rectangle is white. Thus, in this map, a specific color indicates a synteny group. A rectangle is always of the same size as the reference gene in the genome browser; however, it is colored only on part of the gene, which aligns with the compared protein. This allows the user to visualize situations where the alignment is partial. There is one such case in E. coli 536 indicating that the idnK gene in this strain is a pseudogene compared with the idnK gene in CFT073. In contrast with the genome browser, there is no notion of scale on the synteny maps: to see how homologous genes are organized in a synteny group, the user can click on one rectangle in a given synteny group.

ED1a. The foreign origin of this region is also obvious if one looks at the coding prediction curves: the gene model used here does not fit well with the codon usage of the genes annotated in this genomic island. The example shown in Figure 5 also indicates possible paralogy relations through multiple correspondences between genes and one case of frameshift (or sequencing error) in E. coli 536 for the idnK gene (D gluconate kinase; see Figure 5). With such graphical representation, the conservation of genomic context is fully integrated in the process of the expert curation of gene function.

Another visualization mode has been added more recently to represent synteny conservation at different taxonomic levels (i.e. phylum, class, order, family or species). In this 'taxon-synteny' mode (obtained by clicking the 'Switch' button, Figure 5), each line of the synteny map refers to a taxon, and colored boxes represent the percentage of synteny conservation among organisms of the corresponding taxon.

Comparative genomics tools

Computations of homologs and synteny groups between microbial genomes are the starting point of several comparative methods available in the 'Comparative Genomics' menu (Figure 6).

First, the 'Fusion/Fission' functionality provides a list of candidate genes of the selected genome potentially involved in evolutionary events such as gene fusion or fission. Such events involve what is named 'Rosetta-stone' proteins, and suggest a high probability of functional interaction between the involved proteins [23]. Second, the 'Gene phyloprofile' functionality is used to find unique or common genes in the query genome

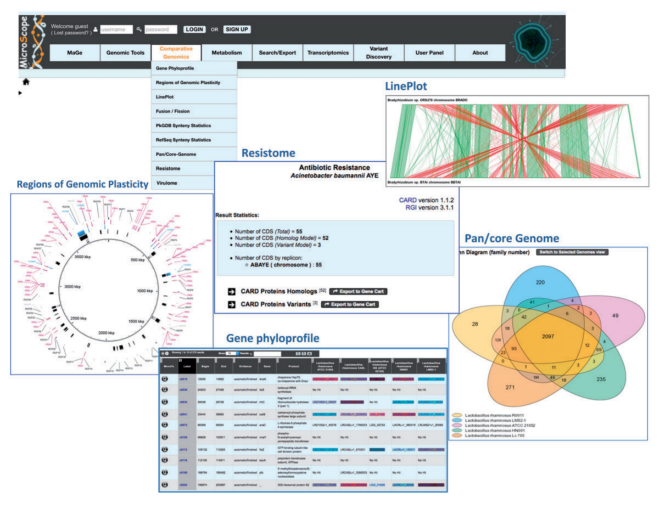


Figure 6. Comparative genomics tools of the MicroScope platform. The figure displays some of the tools available to perform in-depth comparative genomics analyses involving the bacterium of interest and one or a set of organisms: 'Gene Phyloprofile' (comparison of five Lactobacillus rhamnosus strains), 'Line Plot' (shared synteny groups found in the same DNA strand are colored in green, and in red otherwise), 'Regions of Genomic Plasticity' (the predicted genomic island is shown in the second layer of the circular representation), 'Pan-core genome' and 'Resistome'. In this last case, the figure shows Acinetobacter baumannii AYE genes having BLASTP hits with proteins from the CARD database.

with respect to other genomes of interest. Homology constraints and inclusions in synteny group criteria may be applied to refine queries. Third, the 'LinePlot' functionality draws a global graphical representation of conserved syntenies between two selected genomes, and the 'Regions of Genomic Plasticity (RGP)' is used to search for potential horizontal gene transfer (HGT). The method combines (i) the results of algorithms that detect signals in the query sequence indicative of horizontal transfer origin (tRNA hotspots; mobility genes; compositional bias [24]) and (ii) the identification of synteny breaks in the query genome in comparison with closely selected microbial genomes. Results are reported in a tabular form and on a circular representation of the genome (Figure 6). Finally, the 'Pan/ Core Genome' functionality computes dynamically the pangenome and its components (core-genome; variable-genome) of a set of selected organisms (up to 200). The method uses the MicroScope gene families (MICFAM) computed with the SiLiX software [25]. The set of common (= core-genome), variable and strain-specific genes of each compared genomes can be exported in a tabular file format or in a 'Gene Cart'.

Indeed, at any level of the MicroScope Web interface, the gene list that results from the corresponding search/analysis can be selected for inclusion into a 'Gene Cart'. The user can manage several 'Gene Carts' at the same time resulting from different queries. A specific interface has been developed to perform various operations such as the intersection or the difference between two gene carts, to extract sequences or to run multiple alignments via the plugged Jalview software [26] (Functionality 'Gene Carts' of the 'User Panel' menu).

Two functionalities of the 'Comparative Genomics' menu are most specifically related to pathogen analysis (Figure 6): 'Resistome', which uses the Comprehensive Antibiotic Resistance Database [22] a manually curated resource containing high-quality reference data on the molecular basis of antimicrobial resistance, and the Resistance Gene Identifier (RGI) tool to predict the resistome of a genome. The 'Virulome' functionality gives the results of a Blast similarity searches in three distinct data sets of virulence genes: VFDB, which contains experimentally demonstrated virulence genes [27], VirulenceFinder [28] and a subset of the E. coli main virulence genes.

Exploration of metabolic data

The 'Metabolism' menu of MicroScope allows to explore the predicted metabolic pathways using two main resources, KEGG and MetaCyc, and to use analysis tools (Figure 7).

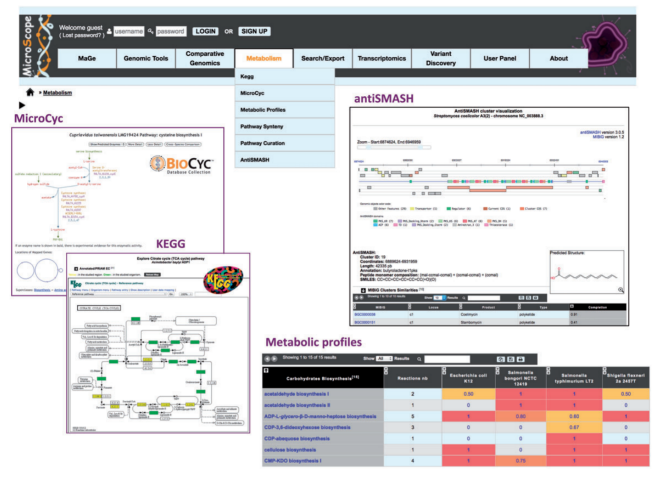


Figure 7. Tools for the analysis of microbial metabolism. Metabolic data can be explored using the KEGG or MetaCyc metabolic pathway hierarchies. On the left, the figure shows, for one selected MicroScope genome, the mapping of the annotated EC numbers on a KEGG metabolic map (enzymes encoded by genes localized on the current genome browser region are highlighted in yellow, and the ones encoded by genes localized elsewhere are highlighted in green). Predicted PGDBs using the Pathway Tools software are available using the 'MicroCyc' functionality. Comparison of metabolic pathways between a set of selected genomes is performed using the 'Metabolic profiles' tool: for each metabolic pathway, a completion value is computed, which corresponds to the number of reactions found in the genome × divided by the total number of reactions in the pathway. This value can take into account pseudogenes or not. It ranges between 0 (absence of the pathway) and 1 (complete pathway). The figure also shows an example of antiSMASH, which predicts Biosynthetic Gene Clusters in prokaryotic genomes. For the NRPS/PKS cluster types, the predicted peptide monomer composition and its corresponding SMILES formula are specified. Below the graphical representation of the predicted antiSMASH cluster, a summary of MIBiG cluster similarities, BGC gene composition as well as tailoring cluster similarities is given.

Starting from the set of predicted and/or validated Enzyme Commission numbers (EC numbers), metabolic maps are dynamically drawn via a request to the KEGG Web server ('KEGG' functionality). A color-based code enables to see the number of enzymatic activities (i.e. EC number) of the annotated genome found in specific metabolic pathways (Figure 7). The interconnected metabolic pathways represented in KEGG are supplemented by the MicroCyc PGDBs built with the Pathway Tools software using MetaCyc as reference metabolic database (see 'Running the annotation pipelines' section). The 'MicroCyc' functionality allows the user to browse and query the metabolic network of a target genome using the Pathway Tools Web interface [18].

These two sets of predicted pathways can be used in the 'Metabolic profiles' functionality. Starting with a selection of organisms and a subset (or all) of metabolic pathways from the KEGG or MetaCyc classification, the tool computes a pathway completion value for each metabolic pathways (Figure 7). These values can be used by the MeV statistical method (Java Web start application) to cluster genomes according to their metabolic capabilities. Moreover, this table is also a good starting point to find candidate genes for missing gene-reaction associations in specific pathways (see example in [6]). In the same way, the 'Pathway Synteny' functionality follows the 'guilt by association' strategy [29], as it combines information on synteny groups and metabolic pathways (i.e. it searches for groups of genes, which share conserved synteny and are found on the same metabolic pathway). Using this interface, annotators can quickly check for reaction-hole candidate genes among the conserved miss-annotated genes of a given group.

Finally, the 'antiSMASH' functionality relies on the integration of the antiSMASH (antibiotics and Secondary Metabolite Analysis Shell) program, which enables rapid genome-wide identification, annotation and analysis of secondary metabolite Biosynthesis Gene Clusters (BGCs) in microbial genomes [30]. Each predicted cluster and its genomic context are explored in a dedicated visualization window showing also a graphical representation of the gene domain composition (Figure 7). For nonribosomal peptide synthetase (NRPS) and polyketide synthase (PKS) cluster types, the predicted peptide monomer composition and its corresponding SMILES formula are specified and the corresponding predicted chemical structure is displayed. For each predicted BGC, a summary of similarities with the

reference database MIBiG [31], BGC gene composition as well as tailoring cluster similarities is given. This last item relies on a knowledge database provided with antiSMASH about tailoring clusters already described in known BGCs and associated with publications.

Analysis of experimental data

The functionalities available in the 'Transcriptomics' and 'Variant discovery' menus rely on the results of the pipelines used to analyze data from transcriptomic projects (i.e. RNA-seq experiments) and data from evolution projects (i.e. clones of the same species at different generation times). Exploration of these experimental data has been illustrated in the two last publications of the MicroScope platform [6, 7].

The 'Transcriptomics' functionality allows exploring the transcript coverage along genome, expression levels of genomic objects (genes, ncRNAs) and differential expression between samples for distinct experimental conditions. All appropriate pairwise comparisons of experimental conditions can be directly queried from the interface. Differentially expressed genes may be projected on reconstructed metabolic networks to highlight metabolic pathways significantly affected by experimental conditions. The 'Variant discovery' functionality offers different tools to explore and analyze the predicted mutations (single nucleotide polymorphisms and small insertions/deletions) in their genomic and functional context. This detection takes into account raw sequencing data and associated read qualities to discriminate between true variations and sequencing errors.

Expert curation of genomic and metabolic data

From the results of the exploration of data and the analysis tools, MicroScope users can review and curate the automatic functional annotation of genes encoded by its genome of interest. This task is performed using the 'Gene Editor', which has been illustrated in the 2013 MicroScope publication [6]. Briefly, it is made of three main sections:

The 'current annotation' section allows the user to modify, delete and add information. The functional description of gene functions is a free-text field exposed to inconsistencies across genes and genomes. We thus have also integrated enumerated lists of well-defined and nonredundant terms for the product type field (defined in GenProtEC [32]), the functional classifications (MultiFun [33] and TIGRFAMs [34]) and for the class field (inspired from the Pseudomonas Genome database [35]), which helps understanding the origin of the functional annotation (e.g. it comes from the functional description of an homologous gene for which the function has been experimentally demonstrated). The curation of associations between genes coding for enzymatic activities and the biochemical reactions catalyzed by these enzymes is performed using two main enzymatic reactions resources: MetaCyc [18] and Rhea [36]. Finally, to alert users about possible inconsistencies, annotation is checked via an automatic procedure launched when the annotation is saved in the database.

The 'automatic annotation' section contains the gene function predicted by our automatic functional annotation procedure ('MicroScope pipeline annotation'), which involves the transfer of the reliable up-to-date reference annotations to 'strong' orthologs, if any [4]. In case of published bacterial genome integrated in MicroScope, the section contains information on the functional annotation in nucleotide sequence databanks and UniProtKB if available.

The 'method results' section provides, for each individual annotation tool executed, a summary of the results, visualized in a tabulated form (this includes precomputed lists of homologs and synteny groups). This integrative strategy allows annotators to quickly browse functional evidences, tracking the history of an annotation and checking the gene context conservation with an orthologous gene having an experimentally demonstrated biological function for example. Criteria for entering an expert annotation are based on different level of evidences from direct experimentation to bioinformatics evidences. The confidence status of each gene annotation is available in the class field of the gene editor. The categories are inspired by the 'protein name confidence' defined in PseudoCAP (Pseudomonas aeruginosa community annotation project). A set of rules allowing to choose this 'class' annotation category according to bioinformatics evidences is proposed in our MicroScope tutorial: https://microscope.readthedocs.io/en/lat est/content/mage/info.html ('How to choose the "Class" annotation category?' and 'Annotation Rules' sections).

Following the integration of novel functionalities into MicroScope, the 'Gene Editor' is constantly evolving. First, new interfaces allowing to ease the curation of resistance and virulence genes are under development, especially using defined ontologies such as ARO, the Antibiotic Resistance Ontology [22]. Second, to fully exploit the results of the different tools dedicated to genomic region analysis (e.g. antiSMASH or RGPfinder), we are currently working on the development of a specific editor to annotate gene clusters such as operons, BGCs, genomic islands, CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats) regions, secretion systems and phages.

Expert annotations are continuously gathered in the MicroScope database. Indeed, ~35 000 annotations are made in a year (Figure 1), and >370 000 genes have been curated so far. A third of these annotations correspond to the description of precise molecular functions supported by direct or indirect (i.e. from homology relationships) experimental evidences. Biologists generally focused their annotations on proteins/functions of interest; however, it is interesting to note that about 50 genomes integrated in MicroScope are near completely curated (≥80% of the genes were expertly annotated), and 124 additional genomes got >300 curated genes.

MicroScope annotations are submitted to INSDC databanks when the genomes get published and can be easily downloaded via the Web interface ('Search/Export->Download Data' functionality). Moreover, we provide a RESTful API to access programmatically public genome data, and semantic Web approaches are currently used to work on the interoperability of MicroScope curated data with other European resources such as UniProtKB [9], HAMAP [37], EnsemblBacteria [1] and Rhea [36]. These developments are performed in the context of the ELIXIR bioinformatics infrastructure (https://www.elixir-europe.org).

Software and database architecture

The technical architecture of the MicroScope platform is shown on Figure 8. Its three components have been described and updated in the previous publications of MicroScope [5, 6]. In summary:

Process management system

The annotation pipelines are organized in a robust automated workflow management system using the jBPM framework (java Business Process Management; http://jbpm.org), which allows

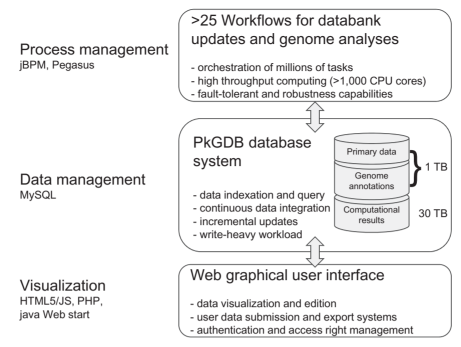


Figure 8. Technical architecture of the MicroScope platform. The MicroScope platform is made of three components: (i) a 'Process management' system to organize workflow execution, (ii) a 'Data management' system, called PkGDB, to store information from databanks, genomes and computational results and (iii) a 'Visualization' system for textual and graphical representation of PkGDB data.

us to handle simultaneously millions of tasks for the analysis of several new microbial genomes. These tasks are parallelized on hundreds of CPU cores using Pegasus MPI cluster module (https://pegasus.isi.edu). The pipelines for the structural, functional and relational annotation orchestrate >50 external/internal bioinformatics software (see section 'Running the MicroScope pipelines'). A large part of these analyses are updated at regular intervals to take into account primary databases growth and new expert annotations.

Data management system

The results of these analysis tools, together with the primary data used as inputs, are stored in a relational database named PkGDB and based on the open-source MySQL relational database management system and the InnoDB (for continuous data integration and incremental updates) and MyISAM (for large bulk inserts) table engines. The PkGDB architecture supports integration of automatic and human-curated functional annotations and records a history of all the modifications. Finally, for metabolic comparative analyses purposes (see the 'Metabolic profiles' functionality in the 'Exploration of metabolic data' section), relational tables have been designed in PkGDB to store information of the MicroCyc PGDBs, together with the KEGG metabolic pathways and modules. The size of PkGDB is today 1 TB for databanks and genome data, and 30 TB for the computational results (Figure 8). Only one instance of the database gathers all genome analyses, which eases collaborative annotation process.

The Web visualization component

The MicroScope Web interface (http://www.genoscope.cns.fr/ agc/microscope) is developed using the Apache/PHP serverbased language and consists of numerous dynamic Web pages containing textual and graphical representations for accessing and querying data. Several useful graphical applications, such as Artemis [38], MeV [39] and IGV [40], are also available in the MicroScope interface through plugged Java applications. As shown in this article, the tools are organized in a menu bar to facilitate the exploration and the curation process. At any level of the interface, a 'Help' functionality is available, and a complete tutorial can be found in the 'About' menu.

Conclusion

In this article, we have described the MicroScope platform from the point of view of the end user, i.e. following one of the main objectives of our prokaryotic genome annotation and comparative system: to allow biologists to submit their genomic data in a simple way and, then, to perform analysis and make relevant assessments of the predicted gene functions using (i) the functionalities for querying and browsing the computed data, (ii) the synteny results and metabolic network predictions, the combination of which can be helpful in formulating hypotheses on the biological function of nonannotated genes and (iii) a gene annotation editor giving access to the results of each method applied, together with links to several useful public resources.

Among the ongoing developments described in the last update of the platform [7], we have currently made great progresses in the consensus representation of thousands bacterial genomes to provide a better analysis workflow of prokaryotic species. The idea is to structure the pan-genome of an organism into the set of 'persistent' genes (relaxed core definition, that is to say genes found in the great majority of the genomes), the 'shell', which gathers moderately conserved genes and the 'cloud' corresponding to rare and unique genes [41]. To organize pangenomic information, we are using a graph data model, where the nodes represent the protein families, and the edges represent the genome co-localization of the two protein families (weighted by the number of the genomes sharing this colocalization). A statistical method is then used to divide the pan-genome into the three main classes (persistent, shell and cloud). The next step is the integration of this representation in MicroScope to facilitate comparative analysis and data visualization of thousands of strains. We will also add functionalities allowing users to select, at any level of this pan-genome graph, a subpart of this graph and, using one genome as reference, to come back to the MaGe genome browser. We are starting to work on an instance of MicroScope based on this novel pangenome representation that will contain most of the reference species found in the human gut microbiota.

Key Points

- MicroScope is open to microbiologists interested in extended analyses of species of interest.
- MicroScope is an integrated environment allowing to perform comparative genomic and metabolic analyses.
- Tools and graphical interfaces for the curation of gene function are part of the specificities of the MicroScope platform.
- MicroScope provides a collaborative environment to share and improve knowledge on genomes.

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